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(54) Title: NUCLEIC ACID SEQUENCES ENCODING A PLANT CYTOPLASMIC PROTEIN INVOLVED IN FATTY ACYL-COA METABOLISM

(57) Abstract

By this invention, a plant β -ketoacyl-CoA synthase condensing enzyme is provided free from intact cells of said plant and capable of catalyzing the production of very long chain fatty acid molecules. Also contemplated are constructs comprising the nucleic acid sequence and a heterologous DNA sequence not naturally associated with the condensing enzyme encoding sequences, and which provide for at least transcription of a plant condensing enzyme encoding sequence in a host cell. In this fashion very long chain fatty acid molecules may be produced in a plant cell. Included are methods of modifying the composition of very long chain fatty acid molecules in a plant cell.

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NUCLEIC ACID SEQUENCES ENCODING A PLANT CYTOPLASMIC PROTEIN INVOLVED IN FATTY ACYL-COA METABOLISM

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This application is a continuation-in-part of of USSN 07/796,256, filed November 20, 1991, a continuation-in-part of USSN 07/933,411, filed August 21, 1992, a continuation-in-part of PCT/US92/09863, filed November 13, 1992, a continuation-in-part USSN 08/066,299, filed May 20, 1993 and a continuation-in-part of USSN 08/160,602, filed November 30, 1993 and a continuation-in-part of of USSN 08/265,047, filed June 23, 1994.

15 Technical Field

The present invention is directed to enzymes, methods to purify, and obtain such enzymes, amino acid and nucleic acid sequences related thereto, and methods of use for such compositions in genetic engineering applications.

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INTRODUCTION

Background

Through the development of plant genetic engineering techniques, it is possible to transform and regenerate a variety of plant species to provide plants which have novel and desirable characteristics. One area of interest for such plant genetic engineering techniques is the production of valuable products in plant tissues. Such applications require the use of various DNA constructs and nucleic acid sequences for use in transformation events to generate plants which produce the desired product. For example, plant functional promoters are required for appropriate expression of gene sequences, such expression being either in the whole plant or in selected plant tissues. addition, selective marker sequences are often used to identify the transformed plant material. Such plant promoters and selectable markers provide valuable tools which are useful in obtaining the novel plants.

One desirable goal, which involves such genetic engineering techniques, is the ability to provide crop plants having a convenient source of wax esters. Wax esters are required in a variety of industrial applications, including pharmaceuticals, cosmetics, detergents, plastics, and lubricants. Such products, especially long chain wax esters, have previously been available from the sperm whale, an endangered species, or more recently, from the desert shrub, jojoba. Neither of these sources provides a convenient supply of way active.

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these sources provides a convenient supply of wax esters. Jojoba is also a plant which synthesizes very long chain fatty acids (VLCFA) in its seed oil. VLCFA are fatty acids having chain lengths longer than 18 carbons. VLCFA are found in the cuticular "waxes" of many plant species as well as in the seed oil of several plant species. Wild type Brassica plants contain VLCFA in their seed oil. Canola is rapeseed that has been bred to eliminate VLCFA from its seed oil. Enzymes involved in the elongation of fatty acids to VLCFA ("elongase" enzymes) have been difficult to characterize at a biochemical level because they are membrane associated (Harwood, JL, "Fatty acid metabolism", Annual rev. of Plant Physiol. and Plant Mol. Biol. (1988) 39:101-38); (von Wettstein-Knowles, PM, "Waxes, cutin, and suberin" in ed. Moore, TS, Lipid Metabolism in Plants (1993), CRC Press, Ann Arbor, pp. 127-166). Although several groups have claimed to partially purify some of these elongase enzymes, to date no one has claimed complete purification of one of these enzymes or cloning of the corresponding genes. von Wettstein-Knowles, PM, (1993) supra; van de Loo, FJ, Fox, BG, and Somerville C. "Unusual fatty acids" in ed. Moore, TS, Lipid Metabolism in Plants, (1993) CRC Press Ann Arbor, pp. 91-

A possible mechanism for fatty acid elongation by the cytoplasmic elongase enzyme system is through a series similar to that found for chloroplast fatty acid synthesis, i.e. via a 4 step reaction (Stumpf and Pollard (1983) supra; van de Loo et al (1993) supra). The first step would be a condensation reaction between malonyl CoA and oleyl

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CoA by ß-ketoacyl-CoA synthase. Then ß-ketoacyl-CoA reductase, ß-hydroxyacyl-CoA dehydratase, and enoyl-CoA reductase ensymes would act sequentially to generate an acyl-CoA molecule elongated by two carbon atoms.

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In order to obtain a reliable source of very long chain fatty acid molecules, such as wax esters or VLCFA, transformation of crop plants, which are easily manipulated in terms of growth, harvest and extraction of products, is desirable. In order to obtain such transformed plants, however, the genes responsible for the biosynthesis of the desired VLCFA or wax ester products must first be obtained.

Wax ester production results from the action of at least two enzymatic activities of fatty acyl CoA metabolism; fatty acyl reductase and fatty acyl:fatty alcohol acyltransferase, or wax synthase. Preliminary studies with such enzymes and extensive analysis and purification of a fatty acyl reductase, indicate that these proteins are associated with membranes, however the enzyme responsible for the fatty acyl:fatty alcohol ligation reaction in wax biosynthesis has not been well characterized. Thus, further study and ultimately, purification of this enzyme is needed so that the gene sequences which encode the enzymatic activity may be obtained.

It is desirable, therefore, to devise a purification protocol whereby the wax synthase protein may be obtained and the amino acid sequence determined and/or antibodies specific for the wax synthase obtained. In this manner, library screening, polymerase chain reaction (PCR) or immunological techniques may be used to identify clones expressing a wax synthase protein. Clones obtained in this manner can be analyzed so that the nucleic acid sequences corresponding to wax synthase activity are identified. The wax synthase nucleic acid sequences may then be utilized in conjunction with fatty acyl reductase proteins, either native to the transgenic host cells or supplied by recombinant techniques, for production of wax esters in host cells.

It would also be desirable to have a gene to an enzyme involved in the formation of very long chain fatty acids. Such a gene could be used to increase the chain length of fatty acids in oilseeds by overexpression of the gene in transgenic plants of virtually any species. The gene could also be used as a probe in low stringency hybridization to isolate homologous clones from other species as a means to clone the gene from other taxa, such as Brassica, Arabidopsis, Crambe, Nasturtium, and Limnanthes, that produce VLCFA. These derived genes could then be used in 10 antisense experiments to reduce the level of VLCFA in the species from which they were isolated, or overexpressed to increase the quantity of VLCFA in transgenic plants of virtually any species. Additionally, the DNA from the homologous Brassica gene encoding this enzyme could be used 15 as a plant breeding tool to develop molecular markers to aid in breeding high erucic acid rapeseed (HEAR) and canola and other oilseed crops. Such techniques would include using the gene itself as a molecular probe or using the DNA 20 sequence to design PCR primers to use PCR based screening techniques in plant breeding programs. Finally, overexpression of the gene in plant epidermal cells could increase cuticle accumulation thereby increasing drought and stress tolerance of transgenic plants over control 25 plants.

Relevant Literature

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Cell-free homogenates from developing jojoba embryos were reported to have acyl-CoA fatty alcohol acyl transferase activity. The activity was associated with a floating wax pad which formed upon differential centrifugation (Pollard et al. (1979) supra; Wu et al. (1981) supra).

Solubilization of a multienzyme complex from Euglena gracilis having fatty acyl-SCoA transacylase activity is reported by Wildner and Hallick (Abstract from The Southwest Consortium Fifth Annual Meeting, April 22-24, 1990, Las Cruces, NM.).

Ten-fold purification of jojoba acyl-CoA: alcohol transacylase protein is reported by Pushnik et al.

(Abstract from The Southwest Consortium Fourth Annual Meeting, February 7, 1989, Riverside, Ca.).

An assay for jojoba acyl-CoA:alcohol transacylase activity was reported by Garver et al. (Analytical Biochemistry (1992) 207:335-340).

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Extracts of developing seeds from HEAR and canola plants were found to differ in their ability to elongate oleyl CoA into VLCFA, with HEAR extracts capable of catalyzing elongation, while canola extracts were not. Stumpf, PK and Pollard MR, "Pathways of fatty acid

10 Stumpf, PK and Pollard MR, "Pathways of fatty acid biosynthesis in higher plants with particular reference to developing rapeseed", in High and Low Erucic Acid Rapeseed Oils (1983) Academic Press Canada, pp. 131-141.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1. The nucleic acid sequence and translated amino acid sequence of a jojoba fatty acyl reductase, as determined from the cDNA sequence, is provided in Figure 1.

- Figure 2. Preliminary nucleic acid sequence and translated amino acid sequence of a jojoba plant cytoplasmic protein involved in fatty acyl-CoA metabolism cDNA clone are provided.
- 10 Figure 3. Nucleic acid and translated amino acid sequences of second class of the jojoba clones, as represented by the sequence of pCGN7614, is provided.
 - Figure 4. Nucleic acid sequence of an oleosin expression cassette is provided.
- 15 Figure 5. Nucleic acid sequence of a *Brassica* condensing enzyme clone, CE15, is provided from a LEAR variety (212).
 - Figure 6. Nucleic acid sequence of a CE20 from the 212 Brassica variety.
- Figure 7. Nucleic acid sequence of a *Brassica* Reston variety (HEAR) clone, of the CE20 class, is provided.
 - Figure 8. Nucleic acid sequence of an Arabadopsis condensing enzyme clone, CE15.
 - Figure 9. Nucleic acid sequence of an Arabadopsis condensing enzyme clone, CE17.

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- Figure 10. Nucleic acid sequence of an Arabadopsis condensing enzyme clone, CE19.
- Figure 11. Partial nucleic acid sequence of Lunaria condensing enzyme clone designated LUN CE8.
- Figure 12. Nucleic acid sequence of a *Lunaria* condensing enzyme clone, Lunaria 1, obtained by probing with LUN CE8.
 - Figure 13. Nucleic acid sequence of a second *Lunaria* condensing enzyme clone obtained from LUN CE8, Lunaria 5.
- Figure 14. Nucleic acid sequence of third *Lunaria* condensing enzyme clone from LUN CE8, Lunaria 27.
 - Figure 15. Nucleic acid sequence to a *Nasturtium* condensing enzyme clone obtained by PCR.

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SUMMARY OF THE INVENTION

By this invention, a DNA sequence encoding a plant cytoplasmic protein involved in fatty acyl-CoA metabolism is provided. Such a sequence is desirable for use in methods aimed at altering the composition of very long chain wax fatty acid related products, such as wax esters and very long chain fatty acids in host cells

In one aspect, the protein of this invention may demonstrate fatty acyl-CoA: fatty alcohol O -acyltransferase activity, such activity being referred to herein as "wax synthase".

In a second aspect, this protein may be required for elongation reactions involved in the formation of very long chain fatty acids. Thus, for example, the protein provides for elongation of C18 fatty acyl CoA molecules to form C20 fatty acids, and also for elongation of C20 fatty acids to form even longer chain fatty acids. It is likely that the elongase activity is the result of B-ketoacyl-CoA synthase activity of this protein, although the possibility exists that the protein provided herein has a regulatory function required for the expression of a S-ketoacyl-CoA synthase or provides one of the other activities known to be involved in acyl-CoA elongation, such as &-ketoacyl-CoA reductase, ß-hydroxyacyl-CoA dehydratase, or enoyl-CoA reductase activities. In any event, the fatty acyl CoA elongation aspect of this protein is referred to herein as "elongase" activity.

The DNA sequence of this invention is exemplified by sequences obtained from a jojoba embryo cDNA library. Several related jojoba sequences have been discovered and are provided in Figures 2 and 3 herein.

In a different aspect of this invention, nucleic acid sequences associated with other proteins related to the exemplified plant cytoplasmic protein involved in fatty

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proteins involved in fatty acyl-CoA metabolism from other plant species, as well as in recombinant constructs for transcription and/or expression in host cells of the protein encoded by such sequences are described. Uses of other nucleic acid sequences associated with the protein encoding sequences are also considered, such as the use of 5' and 3' noncoding regions.

In yet a different aspect of this invention, cells containing recombinant constructs coding for sense and antisense sequences for plant cytoplasmic protein involved in fatty acyl-CoA metabolism are considered. particular, cells which contain the preferred long chain acyl-CoA substrates of the jojoba protein, such as those cells in embryos of Brassica plants, are considered.

In addition, a method of producing a plant cytoplasmic protein involved in fatty acyl-CoA metabolism in a host cell is provided. Accordingly, a plant cytoplasmic protein involved in fatty acyl-CoA metabolism that is recovered as the result of such expression in a host cell is also considered in this invention.

Further, it may be recognized that the sequences of this invention may find application in the production of wax esters in such host cells which contain fatty acyl and fatty alcohol substrates of the wax synthase. Such host cells may exist in nature or be obtained by transformation with nucleic acid constructs which encode a fatty acyl reductase. Fatty acyl reductase, or "reductase", is active in catalyzing the reduction of a fatty acyl group to the corresponding alcohol. Co-pending US patent applications 07/659,975 (filed 2/22/91), 07/767,251 (filed 9/27/91) and 07/920,430 (filed 7/31/92), which are hereby incorporated by reference, are directed to such reductase proteins. This information is also provided in published PCT patent application WO 92/14816. In addition, other sources of wax synthase proteins are described herein which are also desirable sources of reductase proteins. In this regard, plant cells which contain the preferred alcohol substrates of the jojoba wax synthase activity described herein may be prepared by transformation with recombinant nucleic acid

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constructs which encode a fatty acyl reductase nucleic acid sequence.

A further method considered herein involves the production of very long chain fatty acids, or modification of the amounts of such fatty acids, in host cells. Increased production of very long chain fatty acids may be obtained by expression of DNA sequences described herein. On the other hand, antisense constructs containing such sequences may be used to reduce the content of the very long chain fatty acids in a target host organism. In particular, such sense and antisense methods are directed to the modification of fatty acid profiles in plant seed oils and may result in novel plant seed oils having desirable fatty acid compositions.

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DETAILED DESCRIPTION OF THE INVENTION

The nucleic acid sequences of this invention encode a plant cytoplasmic protein involved in fatty acyl-CoA metabolism. Such as a protein includes any sequence of amino acids, such as protein, polypeptide or peptide fragment, which provides the "elongase" activity responsible for production of very long chain fatty acids and for the "wax synthase" activity which provides for esterification of a fatty alcohol by a fatty acyl group to produce a wax ester.

The plant cytoplasmic protein involved in fatty acyl-CoA metabolism of this invention may demonstrate activity towards a variety of acyl substrates, such as fatty acyl-CoA fatty alcohol and fatty acyl-ACP molecules. In addition, both the acyl and alcohol substrates acted upon by the wax synthase may have varying carbon chain lengths and degrees of saturation, although the plant cytoplasmic protein involved in fatty acyl-CoA metabolism may demonstrate preferential activity towards certain molecules.

Many different organisms contain products derived from very long chain fatty acyl-CoA molecules and are desirable sources of a plant cytoplasmic protein involved in fatty acyl-CoA metabolism of this invention. For example, plants

produce epidermal, or cuticular wax (Kolattukudy (1980) in The Biochemistry of Plants (Stumpf, P.K. and Conn, E.E., eds.) Vol.4, p. 571-645), and the desert shrub, jojoba, produces a seed storage wax (Ohlrogge et al. (Lipids (1978) 13:203-210). Such waxes are the result of a wax synthase catalyzed combination of a long chain or very long chain acyl-CoA molecule with a fatty alcohol molecule. Wax synthesis has also been observed in various species of bacteria, such as Acinetobacter (Fixter et al. (1986) J. 10 Gen. Microbiol. 132:3147-3157) and Micrococcus (Lloyd (1987) Microbios 52:29-37), and by the unicellular organism, Euglena (Khan and Kolattukudy (1975) Arch. Biochem. Biophys. 170:400-408). In addition, wax production and wax synthase activity have been reported in microsomal preparations from bovine meibomian glands 15 (Kolattukudy et al. (1986) J. Lipid Res. 27:404-411), avian uropygial glands, and various insect and marine organisms. Consequently, many different wax esters which will have various properties may be produced by wax synthase activity of plant cytoplasmic protein involved in fatty 20 acyl-CoA metabolism of this invention, and the type of wax ester produced may depend upon the available substrate or the substrate specificity of the particular protein of interest.

25 Thus, nucleic acid sequences associated with the plant cytoplasmic protein involved in fatty acyl-CoA metabolism may be cloned into host cells for the production of the enzyme and further studies of the activity. For example, one may clone the nucleic acid encoding sequence into vectors for expression in E. coli cells to provide a ready 30 source of the protein. The protein so produced may also be used to raise antibodies for use in identification and purification of related proteins from various sources, especially from plants. In addition, further study of the protein may lead to site-specific mutagenesis reactions to 35 further characterize and improve its catalytic properties or to alter its fatty alcohol or fatty acyl substrate specificity. A plant cytoplasmic protein involved in fatty acyl-CoA metabolism having such altered substrate

specificity may find application in conjunction with other FAS enzymes.

Prior to the instant invention, amino acid sequences of wax synthase proteins were not known. Thus, in order to obtain the nucleic acid sequences associated with wax synthase, it was necessary to first purify the protein from an available source and determine at least partial amino acid sequence so that appropriate probes useful for isolation of wax synthase nucleic acid sequences could be prepared.

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The desert shrub, Simmondsia chinensis (jojoba) is the source of the encoding sequences exemplified herein. However, related proteins may be identified from other source organisms and the corresponding encoding sequences obtained.

For example, Euglena gracilis produces waxes through the enzymatic actions of a fatty acyl-CoA reductase and a fatty acyl-CoA alcohol transacylase, or wax synthase. Typically, waxes having carbon chain lengths ranging from 24-32 are detected in this organism. The Euglena wax synthase enzyme may be solubilized using a CHAPS/NaCl solution, and a partially purified wax synthase preparation is obtained by Blue A chromatography. In this manner, a 41kD peptide band associated with wax synthase activity is identified.

Acinetobacter species are also known to produce wax ester compositions, although the mechanism is not well defined. As described herein a fatty acyl-CoA alcohol transacylase, or wax synthase activity is detected in Acinetobacter species. The wax synthase activity is solubilized in CHAPS/NaCl, enriched by Blue A column chromatography and may be further purified using such techniques as size exclusion chromatography. By these methods, an approximately 45kD peptide band associated with wax synthase activity is obtained in a partially purified preparation.

In addition, a plant cytoplasmic protein involved in fatty acyl-CoA metabolism which is required for production of very long chain fatty acids may also be found in various

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sources, especially plan sources. In plants, fatty acids up to 18 carbons in chain length are synthesized in the chloroplasts by fatty acid synthase (FAS), a system of several enzymes that elongate fatty acid thioesters of acyl carrier protein (ACP) in 2 carbon increments. reaching the chain length of 18, the thioester linkage is cleaved by a thioesterase, and the fatty acid is transported to the cytoplasm where it is utilized as a coenzyme A (CoA) thioester as acyl-CoA. Further 10 elongation, when it occurs, is catalyzed by an endoplasmic reticulum membrane associated set of elongation enzymes. Very long chain fatty acids (those fatty acids longer than 18 carbons) are found in the cuticular "waxes" of many plant species, and are found in the seed oil of several 15 plant species. The enzymes involved in elongation of fatty acids to VLCFA are membrane associated (Harwood 1988, von Wettstein-Knowles 1993).

Plants which contain desirable "elongase" activities include Arabidopsis, Crambe, Nasturtium and Limnanthes. Thus, the proteins responsible for such elongase activity may be purified and the corresponding encoding sequences identified. Alternatively, such sequences may be obtained by hybridization to the jojoba encoding sequences provided herein.

Although the hydrophobic nature of the proteins of this invention may present challenges to purification, recovery of substantially purified protein can be accomplished using a variety of methods. See, for example, published PCT application WO 93/10241 where purification of jojoba wax synthase protein is described.

Thus, the nucleic acid sequences which encode a plant cytoplasmic protein involved in fatty acyl-CoA metabolism of this invention may be used to provide for transcription of the sequences and/or expression of the protein in host cells, either prokaryotic or eukaryotic.

. Ultimately, stable plant expression in a plant which produces substrates recognized by this enzyme is desired. If a plant targeted for transformation with wax synthase sequences does not naturally contain the fatty alcohol

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and/or fatty acyl ester substrates of this enzyme, a plant extract may be prepared and assayed for activity by adding substrates to the extract. Constructs and methods for transformation of plant hosts are discussed in more detail below.

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As discussed in more detail in the following examples, expression of the nucleic acid sequences provided herein in an initial experiment resulted in increased wax synthase activity. This result, however, was not observed in further *E. coli* expression experiments. In plants, expression of the exemplified sequences (construct pCGN7626, described in Example 8) resulted in production of very long chain fatty acids in a canola type *Brassica*, and modification of the very long chain fatty acid profile in transformed Arabidopsis plants (Example 11).

The nucleic acids of this invention may be genomic or cDNA and may be isolated from cDNA or genomic libraries or directly from isolated plant DNA. Methods of obtaining gene sequences once a protein is purified and/or amino acid sequence of the protein is obtained are known to those skilled in the art.

For example, antibodies may be raised to the isolated protein and used to screen expression libraries, thus identifying clones which are producing the plant 25 cytoplasmic protein involved in fatty acyl-CoA metabolism synthase protein or an antigenic fragment thereof. Alternatively, oligonucleotides may be synthesized from the amino acid sequences and used in isolation of nucleic acid sequences. The oligonucleotides may be useful in PCR to 30 generate a nucleic acid fragment, which may then be used to screen cDNA or genomic libraries. In a different approach, the oligonucleotides may be used directly to analyze Northern or Southern blots in order to identify useful probes and hybridization conditions under which these 35 oligonucleotides may be used to screen cDNA or genomic libraries.

Nucleic acid sequences of this invention include those corresponding to the jojoba plant cytoplasmic protein involved in fatty acyl-CoA metabolism, as well as sequences

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obtainable from the jojoba protein or nucleic acid sequences. By "corresponding" is meant nucleic acid sequences, either DNA or RNA, including those which encode the jojoba plant cytoplasmic protein involved in fatty acyl-CoA metabolism protein or a portion thereof, regulatory sequences found 5' or 3' to said encoding sequences which direct the transcription or transcription and translation (expression) of the protein in jojoba embryos, intron sequences not present in the cDNA, as well as sequences encoding any leader or signal peptide of a precursor protein that may be required for insertion into the endoplasmic reticulum membrane, but is not found in the mature plant cytoplasmic protein involved in fatty acyl-CoA metabolism.

By sequences "obtainable" from the jojoba sequence or protein, is intended any nucleic acid sequences associated with a desired plant cytoplasmic protein involved in fatty acyl-CoA metabolism protein that may be synthesized from the jojoba amino acid sequence, or alternatively identified in a different organism, and isolated using as probes the provided jojoba nucleic acid sequences or antibodies prepared against the jojoba plant cytoplasmic protein involved in fatty acyl-CoA metabolism. In this manner, it can be seen that sequences of these other plant cytoplasmic protein involved in fatty acyl-CoA metabolism may similarly be used to isolate nucleic acid sequences associated with such proteins from additional sources.

For isolation of nucleic acid sequences, cDNA or genomic libraries may be prepared using plasmid or viral vectors and techniques well known to those skilled in the art. Useful nucleic acid hybridization and immunological methods that may be used to screen for the desired sequences are also well known to those in the art and are provided, for example in Maniatis, et al. (Molecular Cloning: A Laboratory Manual, Second Edition (1989) Cold Spring Harbor Laboratory, Cold Spring Harbor, New York).

Typically, a sequence obtainable from the use of nucleic acid probes will show 60-70% sequence identity between the target sequence and the given sequence encoding

a wax synthase enzyme of interest. However, lengthy sequences with as little as 50-60% sequence identity may also be obtained. The nucleic acid probes may be a lengthy fragment of the nucleic acid sequence, or may also be a shorter, oligonucleotide probe. When longer nucleic acid fragments are employed as probes (greater than about 100 bp), one may screen at lower stringencies in order to obtain sequences from the target sample which have 20-50% deviation (i.e., 50-80 sequence homology) from the sequences used as probe. Oligonucleotide probes can be considerably shorter than the entire nucleic acid sequence encoding a wax synthase enzyme, but should be at least about 10, preferably at least about 15, and more preferably at least about 20 nucleotides. A higher degree of sequence identity is desired when shorter regions are used as opposed to longer regions. It may thus be desirable to identify enzyme active sites where amino acid sequence identity is high to design oligonucleotide probes for detecting homologous genes.

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To determine if a related gene may be isolated by hybridization with a given sequence, the sequence is labeled to allow detection, typically using radioactivity, although other methods are available. The labeled probe is added to a hybridization solution, and incubated with filters containing the desired nucleic acids, either Northern or Southern blots (to screen desired sources for homology), or the filters containing cDNA or genomic clones to be screened. Hybridization and washing conditions may be varied to optimize the hybridization of the probe to the sequences of interest. Lower temperatures and higher salt concentrations allow for hybridization of more distantly related sequences (low stringency). If background hybridization is a problem under low stringency conditions, the temperature can be raised either in the hybridization or washing steps and/or salt content lowered to improve detection of the specific hybridizing sequence. Hybridization and washing temperatures can be adjusted based on the estimated melting temperature of the probe as

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discussed in Beltz, et al. (Methods in Enzymology (1983) 100:266-285).

A useful probe and appropriate hybridization and washing conditions having been identified as described above, cDNA or genomic libraries are screened using the labeled sequences and optimized conditions. The libraries are first plated onto a solid agar medium, and the DNA lifted to an appropriate membrane, usually nitrocellulose or nylon filters. These filters are then hybridized with the labeled probe and washed as discussed above to identify clones containing the related sequences.

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For immunological screening, antibodies to the jojoba protein can be prepared by injecting rabbits or mice (or other appropriate small mammals) with the purified protein. Methods of preparing antibodies are well known to those in the art, and companies which specialize in antibody production are also available. Either monoclonal or polyclonal antibodies can be produced, although typically polyclonal antibodies are more useful for gene isolation.

To screen desired plant species, Western analysis is conducted to determine that a related protein is present in a crude extract of the desired plant species, that crossreacts with the antibodies to the jojoba plant cytoplasmic protein involved in fatty acyl-CoA metabolism. This is accomplished by immobilization of the plant extract proteins on a membrane, usually nitrocellulose, following electrophoresis, and incubation with the antibody. Many different systems for detection of the antibody/protein complex on the nitrocellulose filters are available, including radiolabeling of the antibody and second antibody/enzyme conjugate systems. Some of the available systems have been described by Oberfelder (Focus (1989) BRL/Life Technologies, Inc. 11:1-5). If initial experiments fail to detect a related protein, other detection systems and blocking agents may be utilized. When cross-reactivity is observed, genes encoding the related proteins can be isolated by screening expression libraries representing the desired plant species. Expression libraries can be constructed in a variety of

commercially available vectors, including lambda gt11, as described in Maniatis, et al. (supra).

The clones identified as described above using DNA hybridization or immunological screening techniques are then purified and the DNA isolated and analyzed using known techniques. In this manner, it is verified that the clones encode a related protein. Other plant cytoplasmic protein involved in fatty acyl-CoA metabolism may be obtained through the use of the "new" sequences in the same manner as the jojoba sequence was used.

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It will be recognized by one of ordinary skill in the art that nucleic acid sequences of this invention may be modified using standard techniques of site specific mutation or PCR, or modification of the sequence may be accomplished in producing a synthetic nucleic acid sequence. Such modified sequences are also considered in this invention. For example, wobble positions in codons may be changed such that the nucleic acid sequence encodes the same amino acid sequence, or alternatively, codons can be altered such that conservative amino acid substitutions result. In either case, the peptide or protein maintains the desired enzymatic activity and is thus considered part of the instant invention.

A nucleic acid sequence of this invention may be a DNA 25 or RNA sequence, derived from genomic DNA, cDNA, mRNA, or may be synthesized in whole or in part. The gene sequences may be cloned, for example, by isolating genomic DNA from an appropriate source, and amplifying and cloning the sequence of interest using a polymerase chain reaction (PCR). Alternatively, the gene sequences may be 30 synthesized, either completely or in part, especially where it is desirable to provide plant-preferred sequences. Thus, all or a portion of the desired structural gene (that portion of the gene which encodes the protein) may be synthesized using codons preferred by a selected host. 35 Host-preferred codons may be determined, for example, from the codons used most frequently in the proteins expressed in a desired host species.

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The nucleic acid sequences associated with plant cytoplasmic protein involved in fatty acyl-CoA metabolism will find many uses. For example, recombinant constructs can be prepared which can be used as probes or will provide for expression of the protein in host cells. Depending upon the intended use, the constructs may contain the sequence which encodes the entire protein, or a portion thereof. For example, critical regions of the protein, such as an active site may be identified. Further constructs containing only a portion of the sequence which encodes the amino acids necessary for a desired activity may thus be prepared. In addition, antisense constructs for inhibition of expression may be used in which and a portion of the cDNA sequence is transcribed.

Useful systems for expression of the sequences of this invention include prokaryotic cells, such as *E. coli*, yeast cells, and plant cells, both vascular and nonvascular plant cells being desired hosts. In this manner, the plant cytoplasmic protein involved in fatty acyl-CoA metabolism may be produced to allow further studies, such as site-specific mutagenesis of encoding sequences to analyze the effects of specific mutations on reactive properties of the protein.

The DNA sequence encoding a plant cytoplasmic protein involved in fatty acyl-CoA metabolism of this invention may be combined with foreign DNA sequences in a variety of ways. By "foreign" DNA sequences is meant any DNA sequence which is not naturally found joined to the plant cytoplasmic protein involved in fatty acyl-CoA metabolism sequence, including DNA sequences from the same organism which are not naturally found joined to the plant cytoplasmic protein involved in fatty acyl-CoA metabolism sequences. Both sense and antisense constructs utilizing encoding sequences are considered, wherein sense sequence may be used for expression of a plant cytoplasmic protein involved in fatty acyl-CoA metabolism in a host cell, and antisense sequences may be used to decrease the endogenous levels of a protein naturally produced by a target organism. In addition, the gene

sequences of this invention may be employed in a foreign host in conjunction with all or part of the sequences normally associated with the plant cytoplasmic protein involved in fatty acyl-CoA metabolism such as regulatory or membrane targeting sequences.

In its component parts, a DNA sequence encoding a plant cytoplasmic protein involved in fatty acyl-CoA metabolism

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is combined in a recombinant construct having, in the 5' to 3' direction of transcription, a transcription initiation control region capable of promoting transcription and translation in a host cell, the protein encoding sequence and a transcription termination region. Depending upon the host, the regulatory regions will vary, and may include regions from viral, plasmid or chromosomal genes, or the like. For expression in prokaryotic or eukaryotic microorganisms, particularly unicellular hosts, a wide variety of constitutive or regulatable promoters may be employed. Expression in a microorganism can provide a ready source of the plant enzyme. Among transcriptional initiation regions which have been described are regions from bacterial and yeast hosts, such as E. coli, B. subtilis, Sacchromyces cerevisiae, including genes such as beta-galactosidase, T7 polymerase, tryptophan E and the like.

For the most part, the recombinant constructs will involve regulatory regions functional in plants which provide for transcription of the plant cytoplasmic protein involved in fatty acyl-CoA metabolism gene either in the sense or antisense orientation, to produce a functional protein or a complementary RNA respectively. For protein expression, the open reading frame, coding for the plant protein or a functional fragment thereof will be joined at its 5' end to a transcription initiation regulatory region such as the wild-type sequence naturally found 5' upstream to the exemplified jojoba. Numerous other promoter regions from native plant genes are available which provide for a wide variety of constitutive or regulatable, e.g., inducible, expression of structural gene sequences.

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In addition to sequences from native plant genes, other sequences can provide for constitutive gene expression in plants, such as regulatory regions associated with Agrobacterium genes, including regions associated with nopaline synthase (Nos), mannopine synthase (Mas), or octopine synthase (Ocs) genes. Also useful are regions which control expression of viral genes, such as the 35S and 19S regions of cauliflower mosaic virus (CaMV). term constitutive as used herein does not necessarily indicate that a gene is expressed at the same level in all 10 cell types, but that the gene is expressed in a wide range of cell types, although some variation in abundance is often detectable. Other useful transcriptional initiation regions preferentially provide for transcription in certain tissues or under certain growth conditions, such as those 15 from napin, seed or leaf ACP, the small subunit of RUBISCO, and the like.

In embodiments wherein the expression of the plant cytoplasmic protein involved in fatty acyl-CoA metabolism is desired in a plant host, the use of all or part of the complete plant gene may be desired, namely the 5' upstream non-coding regions (promoter) together with the structural gene sequence and 3' downstream non-coding regions may be employed. If a different promoter is desired, such as a promoter native to the plant host of interest or a modified promoter, i.e., having transcription initiation regions derived from one gene source and translation initiation regions derived from a different gene source or enhanced promoters, such as double 35S CaMV promoters, the sequences may be joined together using standard techniques. 30 Additionally, 5' untranslated regions from highly expressed plant genes may be useful to provide for increased expression of the proteins described herein.

The DNA constructs which provide for expression in plants may be employed with a wide variety of plant life, particularly, plants which produce the fatty acyl-CoA substrates of the plant cytoplasmic protein involved in fatty acyl-CoA metabolism, such as Brassica. Other plants of interest produce desirable fatty acyl substrates, such

as medium or long chain fatty acyl molecules, and include but are not limited to rapeseed (Canola varieties), sunflower, safflower, cotton, Cuphea, soybean, peanut, coconut and oil palms, and corn.

As to the fatty alcohol substrate for the ester production, other than jojoba, seed plants are not known to produce large quantities of fatty alcohols, although small amounts of this substrate may be available to the wax synthase enzyme. Therefore, in conjunction with the constructs of this invention, it is desirable to provide the target host cell with the capability to produce fatty alcohols from the fatty acyl molecules present in the host cells. For example, a plant fatty acyl reductase and methods to provide for expression of the reductase enzymes in plant cells are described in co-pending application USSN 07/767,251. The nucleic acid sequence and translated amino acid sequence of the jojoba reductase is provided in Figure Thus, by providing both the wax synthase and reductase activities to the host plant cell, wax esters may be produced from the fatty alcohol and fatty acyl substrates.

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In addition to the jojoba reductase, reductase enzymes from other organisms may be useful in conjunction with the wax synthases of this invention. Other potential sources of reductase enzymes include Euglena, Acinetobacter, Micrococus, certain insects and marine organisms, and specialized mammalian or avian tissues which are known to contain wax esters, such as bovine meibomian glands or avian uropygial glands. Other potential sources of reductase proteins may be identified by their ability to produce fatty alcohols or, if wax synthase is also present, wax esters.

The sequences encoding wax synthase activity and reductase sequences may be provided during the same transformation event, or alternatively, two different transgenic plant lines, one having wax synthase constructs and the other having reductase constructs may be produced by transformation with the various constructs. These plant lines may then be crossed using known plant breeding

techniques to provide wax synthase and reductase containing plants for production of wax ester products.

For applications leading to wax ester production, 5' upstream non-coding regions obtained from genes regulated during seed maturation are desired, especially those preferentially expressed in plant embryo tissue, such as regions derived from ACP, oleosin (Lee and Huang (1991) Plant Physiol. 96:1395-1397) and napin regulatory regions. Transcription initiation regions which provide for preferential expression in seed tissue, i.e., which are undetectable in other plant parts, are considered desirable for wax ester production in order to minimize any disruptive or adverse effects of the gene product in other plant parts. Further, the seeds of such plants may be harvested and the lipid reserves of these seeds recovered 15 to provide a ready source of wax esters. Thus, a novel seed product may be produced in oilseed plants which, absent transformation with wax synthase constructs as described herein, are not known to produce wax esters as a 20 component of their seed lipid reserves.

Similarly, seed promoters are desirable where VLCFA production or inhibition of VLCFA are desired. In this manner, levels of VLCFA may be modulated in various plant species. Such "seed-specific promoters" may be obtained and used in accordance with the teachings of U.S. Serial No. 07/147,781, filed 1/25/88 (now U.S. Serial No. 07/742,834, filed 8/8/81), and U.S. Serial No. 07/494,722 filed on March 16, 1990 having a title "Novel Sequences Preferentially Expressed In Early Seed Development and 30 Methods Related Thereto", all of which co-pending applications are incorporated herein by reference. In addition, where plant genes, such as the jojoba protein is expressed, it may be desirable to use the entire plant gene, including 5' and 3' regulatory regions and any 35 introns that are present in the encoding sequence, for expression of the jojoba genes in a transformed plant species, such as Arabidopsis or Brassica.

Regulatory transcription termination regions may be provided in recombinant constructs of this invention as

well. Transcription termination regions may be provided by the DNA sequence encoding the plant cytoplasmic protein involved in fatty acyl-CoA metabolism or a convenient transcription termination region derived from a different gene source, especially the transcription termination region which is naturally associated with the transcription initiation region. The transcript termination region will contain at least about 0.5kb, preferably about 1-3kb of sequence 3' to the structural gene from which the termination region is derived.

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Additional plant gene regions may be used to optimize expression in plant tissues. For example, 5' untranslated regions of highly expressed genes, such as that of the small subunit (SSU) of RuBP-carboxylase, inserted 5' to DNA encoding sequences may provide for enhanced translation efficiency. Portions of the SSU leader protein encoding region (such as that encoding the first 6 amino acids) may also be used in such constructs. In addition, for applications where targeting to plant plastid organelles is desirable, transit peptide encoding sequences from SSU or other nuclear-encoded chloroplast proteins may be used in conjunction with wax synthase and reductase sequences.

Depending on the method for introducing the DNA expression constructs into the host cell, other DNA sequences may be required. Importantly, this invention is applicable to dicotyledon and monocotyledon species alike and will be readily applicable to new and/or improved transformation and regeneration techniques.

In developing the recombinant construct, the various components of the construct or fragments thereof will normally be inserted into a convenient cloning vector which is capable of replication in a bacterial host, e.g., E. coli. Numerous vectors exist that have been described in the literature. After each cloning, the plasmid may be isolated and subjected to further manipulation, such as restriction, insertion of new fragments, ligation, deletion, insertion, resection, etc., so as to tailor the components of the desired sequence. Once the construct has been completed, it may then be transferred to an

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appropriate vector for further manipulation in accordance with the manner of transformation of the host cell.

Normally, included with the recombinant construct will be a structural gene having the necessary regulatory regions for expression in a host and providing for selection of transformant cells. The gene may provide for resistance to a cytotoxic agent, e.g. antibiotic, heavy metal, toxin, etc., complementation providing prototrophy to an auxotrophic host, viral immunity or the like. Similarly, genes encoding enzymes providing for production

of a compound identifiable by color change, such as GUS, or luminescence, such as luciferase are useful. Depending upon the number of different host species the expression construct or components thereof are introduced, one or more markers may be employed, where different conditions for selection are used for the different hosts.

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In addition to the sequences providing for transcription of sequences encoding the plant cytoplasmic protein involved in fatty acyl-CoA metabolism of this 20 invention, the DNA constructs of this invention may also provide for expression of an additional gene or genes, whose protein product may act in conjunction with the protein described herein to produce a valuable end product. For example, as discussed above, DNA constructs which 25 provide for expression of wax synthase activity and a fatty acyl reductase so that wax esters may produced in transformed hosts, are considered in this invention. Furthermore, production of different wax esters having varying carbon chain lengths and degrees of saturation is 30 desired and may be provided by transforming host plants having fatty alcohol or fatty acyl substrates of varying chain lengths. Such plants may be provided, for example, by methods described in the published international patent application number PCT WO 91/16421, which describes various 35 thioesterase genes and methods of using such genes to produce fatty acyl substrates having varying chain lengths in transformed plant hosts.

Furthermore, to optimize the production of wax esters in oilseed plant hosts, one may wish to decrease the

production of the triacylglyceride oils that are normally produced in the seeds of such plants. One method to accomplish this is to antisense a gene critical to this process, but not necessary for the production of wax esters. Such gene targets include diacylglycerol acyltransferase, and other enzymes which catalyze the synthesis of triacylglycerol. Additionally, it may be desirable to provide the oilseed plants with enzymes which may be used to degrade wax esters as a nutrient source, such as may be isolated from jojoba or various other wax producing organisms. In this manner, maximal production of wax esters in seed plant hosts may be achieved.

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Wax esters produced in the methods described herein may be harvested using techniques for wax extraction from jojoba or by various production methods used to obtain oil products from various oilseed crops. The waxes thus obtained will find application in many industries, including pharmaceuticals, cosmetics, detergents, plastics, and lubricants. Applications will vary depending on the chain length and degree of saturation of the wax ester components. For example, long chain waxes having a double band in each of the carbon chains are liquid at room temperature, whereas waxes having saturated carbon chain components, may be solid at room temperature, especially if the saturated carbon chains are longer carbon chains.

In applications related to elongase activity, the jojoba gene can be used to increase the chain length of fatty acids in oilseeds by overexpression of the gene in transgenic plants of virtually any species; the gene can also be used as a probe in low stringency hybridization to isolate homologous clones from other species that produce VLCFA. These derived genes can then be used in antisense experiments to reduce the level of VLCFA in the species from which they were isolated, or in other plant species where sufficient gene homology is present. Alternatively, these genes could be overexpressed to increase the quantity of VLCFA in transgenic plants.

Additionally, the DNA from the homologous *Brassica* gene encoding this enzyme could be used as a plant breeding

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tool to develop molecular markers to aid in breeding HEAR and canola and other oilseed crops. Such techniques would include using the gene itself as a molecular probe or using the DNA sequence to design PCR primers to use PCR based screening techniques in plant breeding programs.

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Furthermore, overexpression of the gene in plant epidermal cells could increase cuticle accumulation thereby increasing drought and stress tolerance of transgenic plants over control plants.

10 The method of transformation is not critical to the instant invention; various methods of plant transformation are currently available. As newer methods are available to transform crops, they may be directly applied hereunder. For example, many plant species naturally susceptible to 15 Agrobacterium infection may be successfully transformed via tripartite or binary vector methods of Agrobacterium mediated transformation. Other sequences useful in providing for transfer of nucleic acid sequences to host plant cells may be derived from plant pathogenic viruses or 20 plant transposable elements. In addition, techniques of microinjection, DNA particle bombardment, electroporation have been developed which allow for the transformation of various monocot and dicot plant species.

When Agrobacterium is utilized for plant transformation, it may be desirable to have the desired nucleic acid sequences bordered on one or both ends by T-DNA, in particular the left and right border regions, and more particularly, at least the right border region. These border regions may also be useful when other methods of transformation are employed.

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Where Agrobacterium or Rhizogenes sequences are utilized for plant transformation, a vector may be used which may be introduced into an Agrobacterium host for homologous recombination with the T-DNA on the Ti- or Ri-plasmid present in the host. The Ti- or Ri-containing the T-DNA for recombination may be armed (capable of causing gall formation), or disarmed (incapable of causing gall formation), the latter being permissible so long as a functional complement of the vir genes, which encode trans-

acting factors necessary for transfer of DNA to plant host cells, is present in the transformed Agrobacterium host. Using an armed Agrobacterium strain can result in a mixture of normal plant cells, some of which contain the desired nucleic acid sequences, and plant cells capable of gall formation due to the presence of tumor formation genes. Cells containing the desired nucleic acid sequences, but lacking tumor genes can be selected from the mixture such that normal transgenic plants may be obtained.

In a preferred method where Agrobacterium is used as 10 the vehicle for transforming host plant cells, the expression or transcription construct bordered by the T-DNA border region(s) will be inserted into a broad host range vector capable of replication in E. coli and Agrobacterium, there being broad host range vectors described in the 15 literature. Commonly used is pRK2 or derivatives thereof. See, for example, Ditta, et al., (Proc. Nat. Acad. Sci., U.S.A. (1980) 77:7347-7351) and EPA 0 120 515, which are incorporated herein by reference. Alternatively, one may insert the sequences to be expressed in plant cells into a 20 vector containing separate replication sequences, one of which stabilizes the vector in E. coli, and the other in Agrobacterium. See, for example, McBride and Summerfelt (Plant Mol. Biol. (1990) 14:269-276), wherein the pRiHRI (Jouanin, et al., Mol. Gen. Genet. (1985) 201:370-374) 25 origin of replication is utilized and provides for added stability of the plant expression vectors in host Agrobacterium cells.

Utilizing vectors such as those described above, which can replicate in Agrobacterium is preferred. In this 30 manner, recombination of plasmids is not required and the host Agrobacterium vir regions can supply trans-acting factors required for transfer of the T-DNA bordered sequences to plant host cells. For transformation of Brassica cells, Agrobacterium transformation methods may be 35 used. One such method is described, for example, by Radke et al. (Theor. Appl. Genet. (1988) 75:685-694).

The invention now being generally described, it will be more readily understood by reference to the following

examples, which are included for purposes of illustration only and are not intended to limit the invention unless so stated.

EXAMPLES

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Example 1 - Wax synthase Assays

Methods to assay for wax synthase activity in microsomal membrane preparations or solubilized protein preparations are described.

10 Radiolabeled Material

The substrate generally used in the wax synthase assays, [1-14C]palmitoyl-CoA, is purchased from Amersham (Arlington Heights, IL). Other chain length substrates were synthesized in order to perform chain length specification studies. Long chain [1-14C] fatty acids 15 (specific activity 51-56 Ci/mole), namely 11-cis-eicosenoic acid, 13-cis-docosenoic acid and 15-cis-tetracosenoic acid are prepared by the reaction of potassium [14C]cyanide with the corresponding alcohol mesylate, followed by the base 20 hydrolysis of the alcohol nitrile to the free fatty acid. The free fatty acids are converted to their methyl esters with ethereal diazomethane, and purified by preparative silver nitrate thin layer chromatography (TLC). The fatty acid methyl esters are hydrolyzed back to the free fatty acids. Radiochemical purity is assessed by three TLC methods: normal phase silica TLC, silver nitrate TLC, and C18 reversed phase TLC. Radiochemical purity as measured by these methods was 92-98%. Long chain [1-14C] acyl-CoAs are prepared from the corresponding [1-14C] free fatty acids by the method of Young and Lynen (J. Bio. Chem.

(1969) 244:377), to a specific activity of 10Ci/mole. [1-14C]hexadecanal is prepared by the dichromate oxidation of [1-14C]hexadecan-1-ol, according to a micro-scale modification of the method of Pletcher and Tate (Tet. Lett.

35 (1978) 1601-1602). The product is purified by preparative silica TLC, and stored as a hexane solution at -70°C until use.

Assay for Wax synthase Activity in a Microsomal Membrane

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Preparation

Wax synthase activity in a microsomal membrane preparation is measured by incubation of 40µM [1-14C]acyl-CoA (usually palmitoyl-CoA, sp. act. 5.1-5.6 mCi/mmol) and 200µM oleyl alcohol with the sample to be assayed in a total volume of 0.25ml. The incubation mixture also contains 20% w/v glycerol, 1mM DTT, 0.5M NaCl and is buffered with 25mM HEPES (4-[2-hydroxyethyl]-1-piperazineethane-sulfonic acid). HEPES, here and as referred to hereafter is added from a 1M stock solution adjusted to pH 7.5.

A substrate mixture is prepared in a glass vial, with oleyl alcohol being added immediately before use, and is added to samples. Incubation is carried out at 30°C for one hour. The assay is terminated by placing the assay tube on ice and immediately adding 0.25ml isopropanol:acetic acid (4:1 v/v). Unlabeled wax esters (0.1mg) and oleyl alcohol (0.1mg) are added as carriers. The [14C] lipids are extracted by the scaled-down protocol of Hara and Radin (Anal. Biochem. (1978) 90:420). Four ml of hexane/isopropanol (3:2, v/v) is added to the terminated assay. The sample is vortexed, 2ml of aqueous sodium sulphate solution (6.6% w/v) is added, and the sample is again vortexed.

25 C. Assay for Solubilized Wax synthase Activity

For assaying solubilized wax synthase activity, reconstitution of the protein is required. Reconstitution is achieved by the addition of phospholipids (Sigam P-3644, ~40% L-phosphatidyl choline) to the 0.75% CHAPS-solubilized sample at a concentration of 2.5mg/ml, followed by dilution of the detergent to 0.3%, below the CMC. Reconstitution of activity is presumed to be based on the incorporation of wax synthase into the phospholipid vesicles. It is recognized that the amount of wax synthase activity detected after their reconstitution can be influenced by many factors (e.g., the phospholipid to protein ratio and the physical state of the wax synthase protein (e.g. aggregate or dispersed).

D. Analysis of Assay Products

For analyzing the products of either the microsomal membrane preparation wax synthase assay or the solubilized wax synthase assay, two protocols have been developed. One protocol, described below as "extensive assay" is more time-consuming, but yields more highly quantitative results. The other protocol, described below as "quick assay" also provides a measure of wax synthase activity, but is faster, more convenient and less quantitative.

1. Extensive Analysis: Following addition of the sodium sulphate and vortexing the sample, the upper organic phase is removed and the lower aqueous phase is washed with 4ml hexane/isopropanol (7:2 v/v). The organic phases are pooled and evaporated to dryness under nitrogen. The lipid residue is resuspended in a small volume of hexane, and an aliquot is assayed for radioactivity by liquid scintillation counting. The remainder of the sample can be used for TLC analysis of the labeled classes and thereby give a measure of total wax produced.

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For lipid class analysis the sample is applied to a silica TLC plate, and the plate is developed in hexane/diethyl ether/acetic acid (80:20:1 v/v/v). The distribution of radioactivity between the lipid classes, largely wax esters, free fatty acids, fatty alcohols, and polar lipids at the origin, is measured using an AMBIS radioanalytic imaging system (AMBIS Systems Inc., San Diego, CA). If necessary the individual lipid classes can be recovered from the TLC plate for further analysis. Reversed-phase TLC systems using C18 plates developed in methanol have also been used for the analysis.

2. Quick Analysis: Following addition of the sodium sulfate and vortexing the sample, a known percentage of the organic phase is removed and counted via liquid scintillation counting. This calculation is used to estimate the total counts in the organic phase. Another portion of the organic phase is then removed, dryed down under nitrogen, redissolved in hexane and spotted on TLC plates and developed and scanned as described for the detailed assay. In this manner the percentage of the total counts which are incorporated into wax is determined.

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Example 2 - Radiolabeling Wax Synthase Protein

Radiolabeled $[1-1^4C]$ palmitoyl-CoA (Amersham) is added to a wax synthase preparation, either solubilized or a microsomal membrane fraction, in the ratio of 5μ l of label to 40μ l protein sample. The sample is incubated at room temperature for at least 15 minutes prior to further treatment. For SDS-PAGE analysis the sample is treated directly with SDS sample buffer and loaded onto gels for electrophoresis.

Example 3 - Further Studies to Characterize Wax Synthase Activity

15 A. Seed Development and Wax Synthase Activity Profiles
Embryo development was tracked over two summers on
five plants in Davis, CA. Embryo fresh and dry weights
were found to increase at a fairly steady rate from about
day 80 to about day 130. Lipid extractions reveal that
20 when the embryo fresh weight reaches about 300mg (about day
80), the ratio of lipid weight to dry weight reaches the
maximum level of 50%.

Wax synthase activity was measured in developing embryos as described in Example 1. As the jojoba seed coats were determined to be the source of an inhibiting factor(s), the seed coats were removed prior to freezing the embryos in liquid nitrogen for storage at -70° C.

Development profiles for wax synthase activities as measured in either a cell free homogenate or a membrane fraction, indicate a large induction in activity which peaks at approximately 110-115 days after anthesis.

Embryos for enzymology studies were thus harvested between about 90 to 110 days postanthesis, a period when the wax synthase activity is high, lipid deposition has not reached maximum levels, and the seed coat is easily removed. The highest rate of increase of wax synthase activity is seen between days 80 and 90 postanthesis. Embryos for cDNA library construction were thus harvested between about 80 to 90 days postanthesis when presumably the rate of

synthase of wax synthase protein would be maximal. Correspondingly, the level of mRNA encoding wax synthase would be presumed to be maximal at this stage.

B. <u>Substrate Specificity</u>

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Acyl-CoA and alcohol substrates having varying carbon chain lengths and degrees of unsaturation were added to a microsomal membrane fraction having wax synthase activity to determine the range of substrates recognized by the jojoba wax synthase. Wax synthase activity was measured as described in Example 1, with acyl specificity measured using 80μM of acyl-CoA substrate and 100μM of radiolabeled oleyl alcohol. Alcohol specificity was measured using 100μM of alcohol substrate and 40μM of radiolabeled eicosenoyl-CoA. Results of these experiments are presented in Table 1 below.

Table 1

Acyl and Alcohol Substrate Specificity of

Jojoba Wax Synthase

5	Substrate	Wax synthase Activity			
		(pmoles/min)			
	Structure	Acyl Group	Alcohol Group		
	12:0	12	100		
	14:0	95	145		
10	16:0	81	107		
	18:0	51	56		
	20:0	49	21		
	22:0	46	17		
15	18:1	22	110		
	18:2	7	123		
	20:1	122	72		
	22:1	39	41		
	24:1	35	24		
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The above results demonstrate that the jojoba wax synthase utilizes a broad range of fatty acyl-CoA and fatty alcohol substrates.

In addition, wax synthase activity towards various acyl-thioester substrates was similarly tested using palmitoyl-CoA, palmitoyl-ACP and N-acetyl-S-palmitoyl cysteamine as acyl substrates. The greatest activity was observed with the acyl-CoA substrate. Significant activity (~10% of that with acyl-CoA) was observed with acyl-ACP, but no activity was detectable with the N-acetyl-S-palmitoyl cysteamine substrate.

C. <u>Effectors of Activity</u>

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Various sulphydryl agents were screened for their effect on wax synthase activity. Organomercurial compounds were shown to strongly inhibit activity. Iodoacetamide and N-ethylmaleamide were much less effective. Inhibition by para-hydroxymercuribenzoate was observed, but this inhibition could be reversed by subsequent addition of DTT. These results demonstrate that inhibition by para-

hydroxymercuribenzoate involves blocking of an essential sulphydryl group.

D. <u>Size Exclusion Chromatography</u>

A column (1.5cm x 46cm) is packed with Sephacryl-200 (Pharmacia), sizing range: 5,000 - 250,000 daltons) and equilibrated with column buffer (25mM HEPES, 20% glycerol, 0.75% CHAPS, 1mM EDTA) containing 0.5M NaCl. Approximately 2 ml of a pooled concentrate from a single 1.5 M NaCl elution from a Blue A column (see Ex. 4C) is loaded and the column run at 0.5 ml/min. The eluted fractions are assayed for wax synthase activity according to the reconstitution protocol described in Example 1. Wax synthase activity appears as a broad peak beginning at the void fraction and decreasing throughout the remainder of the run. A portion of the fractions having wax synthase activity are treated 15 with 1-14C 16:0-CoA (0.0178 uM) for 15 minutes at room temperature. SDS is added to 2% and the samples are loaded on an SDS-PAGE gel. Following electrophoresis, the gel is blotted to Problott (Applied Biosystems; Foster City, CA) 20 and the dried blot membrane analyzed by autoradiography. Alternatively, the blot may be scanned for radioactivity using an automated scanning system (AMBIS; San Diego, Ca.). In this manner, it is observed that the 57kD radiolabeled band tracks with wax synthase activity in the analyzed 25 fractions.

Protein associated with wax synthase activity is further characterized by chromatography on a second size exclusion matrix. A fraction (100ul) of a 10% concentrated 1.5M NaCl elution from a Blue A column (following a 1.0M NaCl elution step) which contains wax synthase activity is chromatographed on a Superose 12 HR10/30 column (Pharmacia: Piscataway, NJ) and analyzed by Fast Protein Liquid Chromatography (FPLC) on a column calibrated with molecular weight standards (MW GF-70 and MW GF-1000; Sigma).

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Activity assays are performed on the eluted fractions.

Most 53% of the recovered wax synthase activity is found in
the void fractions, but an easily detectable activity is
found to elute at ~55kd according to the calibration curve.

These data indicate the minimum size of an active native

wax synthase protein is very similar to the 57kD size of the labeled band, thus providing evidence that wax synthase activity is provided by a single polypeptide. The fraction of wax synthase activity observed in the void fractions is presumably an aggregated form of the enzyme.

E. Palmitoyl-CoA Agarose Chromatography

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A column (1.0 x 3cm) is packed with 16:0-CoA agarose (Sigma P-5297) and equilibrated with column buffer (See, Example 1, D.) containing 0.2M NaCl. Approximately 4 ml of a pooled concentrate from the 1.5M NaCl wash of the Blue A column is thawed and the salt concentration reduced by passage of the concentrate over a PD-10 (Pharmacia) desalting column equilibrated in 0.2M NaCl column buffer. The reduced salt sample (5ml) is loaded onto the 16:0 CoA agarose column at a flow rate of 0.15 ml/min. The column is washed with 0.5M NaCl column buffer and then with 1.5M NaCl column buffer. Although some wax synthase activity flows through the column or is removed by the 0.5M NaCl wash, the majority of the recovered activity (21% of the loaded activity) is recovered in the 1.5M NaCl eluted peak.

Portions of the fractions which demonstrate wax synthase activity are radiolabeled with [14C]palmitoyl-CoA as described in Example 2 and analyzed by SDS polyacrylamide gel electrophoresis (Laemmli, Nature (1970) 227:680-685). Again the approximate 57kD radio labelled protein band is observed to track with wax synthase activity.

Example 4 - Purification of Jojoba Wax Synthase

Methods are described which may be used for isolation of a jojoba membrane preparation having wax synthase activity, solubilization of wax synthase activity and further purification of the wax synthase protein.

A. <u>Microsomal Membrane Preparation</u>

Jojoba embryos are harvested at approximately 90-110 days after flowering, as estimated by measuring water content of the embryos (45-70%). The outer shells and seed coats are removed and the cotyledons quickly frozen in liquid nitrogen and stored at -70°C for future use. For

initial protein preparation, frozen embryos are powdered by pounding in a steel mortar and pestle at liquid nitrogen temperature. In a typical experiment, 70g of embryos are processed.

- The powder is added, at a ratio of 280ml of solution per 70g of embryos, to the following high salt solution: 3M NaCl, 0.3M sucrose, 100mM HEPES, 2mM DTT, and the protease inhibitors, 1mM EDTA, 0.7µg/ml leupeptin, 0.5µg/ml pepstatin and 17µg/ml PMSF. A cell free homogenate (CFH)
- is formed by dispersing the powdered embryos in the buffer with a tissue homogenizer (Kinematica, Switzerland; model PT10/35) for approximately 30 sec. and then filtering through three layers of Miracloth (CalBioChem, LaJolla, CA). The filtrate is centrifuged at 100,000 x g for one hour.

The resulting sample consists of a pellet, supernatant and a floating fat pad. The fat pad is removed and the supernatant fraction is collected and dialyzed overnight (with three changes of the buffering solution) versus a solution containing 1M NaCl, 100mM HEPES, 2mM DTT and 0.5M EDTA. The dialyzate is centrifuged at 200,000 x g for 1 1/2 hour to yield a pellet, DP2. The pellet is suspended in 25mM HEPES and 10% glycerol, at 1/20 of the original CFH volume, to yield the microsomal membrane preparation.

Activity is assayed as described in Example 1.

Recovery of wax synthase activity is estimated at 34% of the original activity in the cell free homogenate. Wax synthase activity in this preparation is stable when stored at -70°C.

Solubilization of Wax synthase Protein

CHAPS (3-[(3-cholamidopropyl)-dimethyl-ammonio]-1propanesulfonate) and NaCl are added to the microsomal membrane preparation to yield final concentrations of 2% and 0.5M, respectively. The samples are incubated on ice for approximately one hour and then diluted with 25 mMHEPES, 20% glycerol, 0.5M NaCl to lower the CHAPS concentration to 0.75%. The sample is then centrifuged at $200,000 \ \mathrm{x}$ g for one hour and the supernatant recovered and assayed for wax synthase activity as described in Example Typically, 11% of the wax synthase activity from the microsomal membrane preparation is recovered in the supernatant fraction. The solubilized wax synthase activity is stable when stored at -70°C.

Blue A Column Chromatography 15 C.

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A column (2.5 \times 8cm) with a bed volume of approximately 30ml is prepared which contains Blue A (Cibacron Blue F3GA; Amicon Division, W.R. Grace & Co.), and the column is equilibrated with the column buffer (25 mMHEPES, 20% glycerol, 0.75% CHAPS, 1mM EDTA) containing 0.4M NaCl. The solubilized wax synthase preparation is diluted to 0.4M NaCl by addition of column buffer (25mM HEPES, 20% glycerol, 0.75% CHAPS, 1mM EDTA) and loaded to the Blue A column.

25 The column is washed with column buffer containing 0.5M NaCl until no protein can be detected (as measured by absorbance at 280nm) in the buffer flowing through the column. Greater than 94% of the wax synthase activity binds to the column, while greater than 83% of other protein passes through. Typically, approximately 20% of the loaded 30 wax synthase activity is recovered by elution. A portion of the recovered activity (17%) elutes with a 1.0M NaCl column buffer wash, while approximately 75% of the recovered activity elutes as a broad peak in a 150ml wash with 1.5M NaCl column buffer. Five ml fractions of the 1.5M wash are collected and assayed for wax synthase activity as described in Example 1. Fractions containing wax synthase activity are pooled and concentrated ten fold using an Amicon stirred cell unit and a YM30 membrane. The

concentrated wax synthase preparation may be stored at -70°C.

D. <u>Size Exclusion Column Chromatography</u>

In fractions collected from chromatography on Blue A
the acyl-transferase enzyme activity responsible for
formation of wax esters from fatty alcohol and acyl-CoA coelutes with the measurable activity of ß-ketoacyl-CoA
synthase. The ß-ketoacyl-CoA synthase activity can be
separated from this wax synthase activity through size
exclusion chromatography using S 100 sepharose. The
preferred column buffer for size exclusion chromatography
comprises 1.0% CHAPS, as at 0.75% CHAPS the enzyme tends to
aggregate, i.e., stick to itself and other proteins. Using
a column buffer adjusted to 1.0% CHAPS allows clean
separation of the activity of wax synthase on S 100, wax
synthase being retained, from the ß-ketoacyl-CoA synthase
protein, the latter being voided. The majority of wax
synthase activity elutes from the S 100 sizing column as a

peak with a molecular mass ~ of 57 kDa. At 0.75% CHAPS

20 only a small portion of total assayable wax synthase
activity is found at 57 kDa, with the remainder distributed
over void and retained fractioins.

Wax synthase also has an estimated molecular mass of ~57 kDa based on SDS gels of radiolabelled protein, i.e., wax synthase protein which has been labeled by the procedure described above by incubation with 14C-palmitoyl-CoA. The labelled band tracks with wax synthase activity in fractions collected from a size exclusion column, while ß-ketoacyl-CoA synthase activity is completely voided by the S 100 column.

As a predominant 57 kDa protein from the Blue A column fraction, the ß-ketoacyl-CoA synthase can be amino acid sequenced from bands removed from SDS PAGE. Wax synthase activity can be isolated by SDS PAGE and cloned by a similar procedure from fractions retained on S 100.

E. SDS PAGE Analysis

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Samples from the S 100 or active BlueA column fractions are diluted in SDS PAGE sample buffer (1x buffer = 2% SDS, 30mM DTT, 0.001% bromphenol blue) and analyzed by

electrophoresis on 12% tris/glycine precast gels from NOVEX (San Diego, CA). Gels are run at 150V, constant voltage for approximately 1.5 hours. Protein is detected by silver staining (Blum et al., Electrophoresis (1987) 8:93-99). Careful examination of the gel reveals only a few polypeptides, including one of approximately 57kD, whose staining intensity in the various fractions can be correlated with the amount of wax synthase activity detected in those fractions. Furthermore, if radiolabeled [1-14C] palmitoyl-CoA is added to the protein preparation prior to SDS PAGE analysis, autoradiography of the gel reveals that the 57kD labeled band tracks with wax synthase activity in these fractions. Other proteins are also present in the preparation, including the 56 and 54kD reductase proteins described in co-pending application USSN 07/767,251.

Continuous Phase Elution

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Wax synthase protein is isolated for amino acid sequencing using an SDS-PAGE apparatus, Model 491 Prep Cell (Bio-Rad Laboratories, Inc., Richmond, CA), according to 20 manufacturer's instructions. A portion (15 ml) of the wax synthase activity from the 1.5M NaCl elution of the Blue A column is concentrated 10 fold in a Centricon 30 (Amicon Division, W. R. Grace & Co.; Beverly, MA) and desalted with column buffer on a Pharmacia PD-10 desalting column. sample is treated with 2% SDS and a small amount of bromphenol blue tracking dye and loaded onto a 5 ml, 4% acrylamide stacking gel over a 20 ml, 12% acrylamide running gel in the Prep Cell apparatus. The sample is electrophoresed at 10W and protein is continuously 30 collected by the Prep Cell as it elutes from the gel. eluted protein is then collected in 7.5-10 ml fractions by a fraction collector. One milliliter of each fraction in the area of interest (based on the estimated 57kD size of the wax synthase protein) is concentrated to 40 μl in a 35 Centricon 30 and treated with 2% SDS. The samples are run on 12% acrylamide mini-gels (Novex) and stained with Various modifications to the continuous phase elution process in order to optimize for wax synthase

recovery may be useful. Such modifications include adjustments of acrylamide percentages in gels volume of the gels, and adjustments to the amount of wax synthase applied to the gels. For example, to isolate greater amounts of the wax synthase protein the Blue A column fractions may be applied to larger volume, 20-55 ml, acrylamide gels at a concentration of approximately 1 mg of protein per 20 ml of gel. The protein fractions eluted from such gels may then be applied 10-15% gradient acrylamide gels for increased band separation.

The protein content of each fraction is evaluated visually and fractions containing wax synthase protein are pooled and concentrated for amino acid sequencing. order to maximize the amount of wax synthase enzyme collected, fractions which also contain the 56kD reductase protein band are included in the pooled preparation. As the reductase protein sequence is known (see Figure 1), further purification of wax synthase protein in the pooled preparation is not necessary prior to application of amino 20 acid sequencing techniques (see Example 5).

Blotting Proteins to Membranes

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Alternatively, wax synthase protein may be further isolated for amino acid sequencing by transfer to PVDF membranes following SDS-PAGE, either Immobilon-P 25 (Millipore; Bedford, MA) or ProBlott (Applied Biosystems; Foster City, CA). Although transfer to nitrocellulose may also be useful, initial studies indicate poor transfer to nitrocellulose membranes, most likely due to the hydrophobic nature of this protein. PVDF membranes, such as ProBlott and Immobilon-P find preferential use in 30 different methods, depending on the amino acid sequencing technique to be employed. For example, transfer to ProBlott is useful for N-terminal sequencing methods and for generation of peptides from cyanogen bromide digestion, Immobilon-P is preferred. 35

Blotting to Nitrocellulose: When protein is electroblotted to nitrocellulose, the blotting time is typically 1-5 hours in a buffer such as 25mM Tris, 192mM glycine in 5-20% methanol. Following electroblotting,

membranes are stained in 0.1% (w/v) Ponceau S in 1% (v/v) acetic acid for 2 minutes and destained in 2-3 changes of 0.1% (v/v) acetic acid, 2 minutes for each change. These membranes are then stored wet in heat-sealed plastic bags at -20° C. If time permits, blots are not frozen but used immediately for digestion to create peptides for determination of amino acid sequence as described below.

Blotting to PVDF: When protein is electroblotted to Immobilon P PVDF, the blotting time is generally about 10 1-2 hours in a buffer such as 25mM Tris/192mM glycine in 20% (v/v) methanol. Following electroblotting to PVDF, membranes are stained in 0.1% (w/v) Coomassie Blue in 50% (v/v) methanol/10% (v/v) acetic acid for 5 minutes and destained in 2-3 changes of 50% (v/v) methanol/10% (v/v)acetic acid, 2 minutes for each change. PVDF membranes are 15 then allowed to air dry for 30 minutes and are then stored dry in heat-sealed plastic bags at -20°C. Protein blotted to PVDF membranes such as Pro Blott, may be used directly to determine N-terminal sequence of the intact protein. A protocol for electroblotting proteins to ProBlott is 20 described below in Example 5A.

Example 5 - Determination of Amino Acid Sequence

In this example, methods for determination of amino acid sequences of plant proteins associated with wax synthase activity are described.

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A. <u>Cyanogen Bromide Cleavage of Protein and Separation of Peptides</u>

Cyanogen bromide cleavage is performed on the protein of interest using the methodology described in the Probe-Design Peptide Separation System Technical Manual from Promega, Inc. (Madison, WI). The wax synthase protein, if not available in a purified liquid sample, is blotted to a PVDF membrane as described above. Purified wax synthase protein or wax synthase bands from the PVDF blot, are placed in a solution of cyanogen bromide in 70% (v/v) formic acid, and incubated overnight at room temperature. Following this incubation the cyanogen bromide solutions are removed, pooled and dried under a continuous nitrogen

stream using a Reacti-Vap Evaporator (Pierce, Rockford, IL). Additional elution of cyanogen bromide peptides from PVDF may be conducted to ensure complete removal, using a peptide elution solvent such as 70% (v/v) isopropanol, 0.2% (v/v) trifuoroacetic acid, 0.1mM lysine, and 0.1mM thioglycolic acid. The elution solvents are then removed and added to the tube containing the dried cyanogen bromide solution, and dried as described above. The elution procedure may be repeated with fresh elution solvent. $50\mu l$ of HPLC grade water is then added to the dried peptides and the water removed by evaporation in a Speed-Vac (Savant, Inc., Farmingdale, NY).

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Peptides generated by cyanogen bromide cleavage are separated using a Tris/Tricine SDS-PAGE system similar to that described by Schägger and von Jagow (Anal. Biochem. 15 (1987) 166:368-379). Gels are run at a constant voltage of 125-150 volts for approximately 1 hour or until the tracking dye has begun to run off the bottom edge of the gel. Gels are soaked in transfer buffer (125mM Tris, 50mM glycine, 10% (v/v) methanol) for 15-30 minutes prior to 20 transfer. Gels are blotted to ProBlott sequencing membranes (Applied Biosystems, Foster City, CA) for 2 hours at a constant voltage of 50 volts. The membranes are stained with Coomassie blue (0.1% in 50% (v/v) methanol/10% (v/v) acetic acid) and destained for 3X 2 min. in 50% (v/v)methanol/10% (v/v) acetic acid. Membranes are air-dried for 30-45 minutes before storing dry at -20° C.

Peptides blotted on to ProBlott can be directly loaded to the sequencer cartridge of the protein sequencer without the addition of a Polybrene-coated glass fibre filter. Peptides are sequenced using a slightly modified reaction cycle, BLOT-1, supplied by Applied Biosystems. Also, solution S3 (butyl chloride), is replaced by a 50:50 mix of S1 and S2 (n-heptane and ethyl acetate). These two modifications are used whenever samples blotted to ProBlott are sequenced.

B. <u>Protease Digestion and Separation of Peptides</u>

Purified wax synthase protein provided in a liquid

solution or wax synthase proteins blotted to nitrocellulose

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may be subjected to digestion with proteases in order to obtain peptides for sequencing. The method used is that of Aebersold, et al. (PNAS (1987) 84:6970).

For protein provided on nitrocellulose, bands of the wax synthase proteins, and also an equal amount of blank nitrocellulose to be used as a control, are cut out of the nitrocellulose membrane and washed several times with HPLC grade water in order to remove the Ponceau S. Following this wash, 1.0ml of 0.5% polyvinylpyrrolidone (PVP-40,

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Aldrich, Milwaukee, WI) in 0.5% acetic acid is added to the membrane pieces and this mixture is incubated for 30 minutes at 37°C. In order to remove the PVP-40 completely, nitrocellulose pieces are washed with many volumes of HPLC grade water (8 x 5ml), checking the absorbance of the washes at 214nm on a spectrophotometer. Also, PVP-40 is more easily removed if bands are not cut into small pieces until after PVP-40 treatment and washing.

The proteins, in solution or on nitrocellulose pieces, are then suspended in an appropriate digest buffer, for example trypsin digest buffer, 100mM sodium bicarbonate pH 8.2, or endoproteinase gluC buffer, 25mM ammonium carbonate/1mM EDTA, pH 7.8. Acetonitrile is added to the digest mixture to a concentration of 5-10% (v/v). Proteases are diluted in digest buffer and added to the

digest mixture, typically at a ratio of 1:10 (w/w) protease to protein. Digests are incubated 18-24 hours. For example, trypsin digests are incubated at 37°C and endoproteinase gluC digests are incubated at room temperature. Similarly, other proteases may be used to digest the wax synthase proteins, including lysC and aspN. While the individual digest buffer conditions may be different, the protocols for digestion, peptide separation,

purification and sequencing are substantially the same as those described for digestion with trypsin and gluC.

Following overnight incubation, digest reactions are stopped by the addition of 10 μ l 10% (v/v) trifluoroacetic acid (TFA) or 1 μ l 100% TFA. When the protein is provided on nitrocellulose, the nitrocellulose pieces are washed with 1-5 100 μ l volumes of digest buffer with 5-10%

acetonitrile, and these volumes are concentrated to a volume of less than 100 μl in a Speed-Vac.

The peptides resulting from digestion are separated on a Vydac reverse phase C18 column (2.1mm x 100mm) installed in an Applied Biosystems (Foster City, CA) Model 130 High Performance Liquid Chromatograph (HPLC). Mobile phases used to elute peptides are: Buffer A: 0.1mM sodium phosphate, pH2.2; Buffer B: 70% acetonitrile in 0.1mM sodium phosphate, pH2.2. A 3-step gradient of 10-55% buffer B over two hours, 55-75% buffer B over 5 minutes, and 75% buffer B isocratic for 15 minutes at a flow rate of 50µl/minute is used. Peptides are detected at 214nm, collected by hand, and then stored at -20° C.

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Due to the hydrophobic nature of the wax synthase 15 proteins, addition of a detergent in enzyme digestions buffers may be useful. For example, fractions from the continuous phase elution procedure described above which contain the jojoba wax synthase are concentrated in a Centricon 30 in 100mM NaHCO3/1.0% CHAPS to a final volume of 110 μ l. Two μ g of trypsin in 5 μ l of 100mM Na HCO3/1.0% CHAPS is added to the protein solution and the mixture is incubated overnight at 37°C, and the digestion stopped by addition of trifluoroacetic acid (TFA). The sample is centrifuged lightly and the peptides separated on a Vydac C18 column and eluted as described above. In this 25 procedure, the CHAPS elutes at $\sim40-53\%$ Buffer B, and obscures the peptide peaks in this region.

Where the primary separation yields a complex peptide pattern, such as where excess protein is used or

contaminants (such as the jojoba reductase protein) are present, peptide peaks may be further chromatographed using the same column, but a different gradient system. For the above jojoba wax synthase preparation, hydrophilic peaks were separated using a gradient of 0-40% Buffer B for 60 minutes, 40-75% B for 35 minutes and 75-100% B for 10 minutes. Hydrophobic peaks were separated using 0-40% Buffer B for 40 minutes, 40-80% B for 60 minutes and 80-100% B for 10 minutes. For these separations, Buffer A is 0.1% TFA and Buffer B is 0.1% TFA in acetonitrile.

C. N-terminal Sequencing of Proteins and Peptides

All sequencing is performed by Edman degradation on an Applied Biosystems 477A Pulsed-Liquid Phase Protein Sequencer; phenylthiohydantoin (PTH) amino acids produced by the sequencer are analyzed by an on-line Applied Biosystems 120A PTH Analyzer. Data are collected and stored using an Applied Biosystems model 610A data analysis system for the Apple Macintosh and also on to a Digital Microvax using ACCESS*CHROM software from PE NELSON, Inc. (Cupertino, CA). Sequence data is read from a chart recorder, which receives input from the PTH Analyzer, and is confirmed using quantitative data obtained from the

recorder, which receives input from the PTH Analyzer, are is confirmed using quantitative data obtained from the model 610A software. All sequence data is read independently by two operators with the aid of the data analysis system.

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For peptide samples obtained as peaks off of an HPLC, the sample is loaded on to a Polybrene coated glass fiber filter (Applied Biosystems, Foster City, CA) which has been subjected to 3 pre-cycles in the sequencer. For peptides which have been reduced and alkylated, a portion of the PTH-amino acid product material from each sequencer cycle is counted in a liquid scintillation counter. For protein samples which have been electroblotted to Immobilon-P, the band of interest is cut out and then placed above a Polybrene coated glass fiber filter, pre-cycled as above and the reaction cartridge is assembled according to manufacturer's specifications. For protein samples which have been electroblotted to ProBlott, the glass fiber filter is not required.

In order to obtain protein sequences from small amounts of sample (5-30 pmoles), the 477A conversion cycle and the 120A analyzer as described by Tempst and Riviere (Anal. Biochem. (1989) 183:290).

Amino acid sequence of jojoba peptides obtained by 35 trypsin digestion as described above are presented in Table 2 below.

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Table 2

Amino Acid Sequence of Jojoba 57 kDa protein Tryptic

Peptides

J		
	SQ1114	ETYVPESVTKK
	SQ1084	VPXEPSIAAX
	SQ1083	ETYVPEEvtk
	SQ1120	DLMAVAGEA1k
10	SQ1125	MTNVKPYIPDF
	SQ1129	FLPXXVAiTGe
	SQ1131	FGNTSSXXLyxelayak
	SQ1137	AEAEEVMYGAIDEVLEK

- The amino acid sequence is represented using the one letter code. "X" represents a position where the amino acid could not be identified, and amino acids represented by lower case letters represent residues which were identified with a lesser degree of confidence.
- 20 Example 6 ~ Purification of Additional Wax Synthases

and Reductases

A. Adaptation of jojoba wax synthase solubilization and purification methods to obtain partially purified preparations of wax synthase from other organisms are described.

Acinetobacter

Cells of Acinetobacter calcoaceticus strain BD413
(ATCC #33305) are grown on ECLB (E. coli luria broth),

collected during the logarithmic growth phase and washed in a buffer containing; Hepes, pH 7.5, 0.1M NaCl, 1mM DTT and protease inhibitors. Washed cells were resuspended in fresh buffer and ruptured by passage through a French pressure cell (two passes at ~16,000p.s.i.). Unbroken

cells are removed by centrifugation at 5000 x g for 10 minutes, and membranes are collected by centrifugation at 100,000 x g for 1 hour. The membrane pellet is homogenized in storage buffer (25mM Hepes, pH 7.5, 10% (w/v) glycerol). Wax synthase activity is detected in these membranes using

assay conditions described for the jojoba enzyme in Example

1B, using [1-14C] palmitoyl-CoA and 18:1 alcohol as the substrates.

Wax synthase activity is solubilized by incubation of the membranes with 2% CHAPS in the presence of 0.5M NaCl, as described for the jojoba enzyme in Example 4B. Solubilization of the activity is demonstrated by the detection of wax synthase enzyme activity in the supernatant fraction after centrifugation at 200,000g for 1 hour and by size exclusion chromatography (i.e. the 10 activity elutes from the column in the retained fractions as a symmetrical peak). The activity of the solubilized enzyme is detected by simple dilution of the CHAPS concentration to ~0.3% (i.e. to below its CMC). 15

Incorporation of the enzyme into phospholipid vesicles is not required to detect solubilized activity.

For purification, the solubilized Acinetobacter wax synthase activity is subjected to chromatographic purification procedures similar to those described for the jojoba acyl-CoA reductase. The soluble protein preparation is loaded to a Blue A agarose column under low salt conditions (150mM NaCl in a column buffer containing 0.75% CHAPS, 10% glycerol, 25mM Hepes, pH 7.5) and eluted from the column using 1.0M NaCl in the column buffer.

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Size exclusion chromatography on Superose 12 (Pharmacia; Piscataway, NJ) medium is used to obtain an estimate of the size of the native enzyme and to aid in identifying candidate polypeptides. Comparison to molecular mass standards chromatographed under identical conditions yields an estimate of ~46kD for the native wax synthase activity. Three polypeptides bands, with apparent molecular masses of 45kD, 58kD and 64kD, were identified which tracked with wax synthase activity. N-terminal sequence of the 45kD polypeptide, the strongest candidate for wax synthase, is determined as XDIAIIGSGsAGLAQaxilkdag, where the one letter code for amino acids is used, "X" represents a position where the amino acid could not be identified, and amino acids represented by lower case letters represent residues which were identified with a

lesser degree of confidence. In addition, sequence of a tryptic peptide of the *Acinetobacter* wax synthase protein is determined as QQFTVWXNASEPS.

Euglena

5 Euglena gracilis, strain Z (ATCC No. 12716) is grown heterotrophically in the dark (Tani et al. (1987) Agric. Biol. Chem. 51:225-230) at ~26°C with moderate shaking. Cells are collected and washed in buffer containing 25mMBis-Tris-Propane, pH 7.0, 0.25M NaCl and 1mM EDTA. Washed cells are resuspended in fresh buffer and ruptured by passage through a French pressure cell (two passes at ~16,000 p.s.i.). Unbroken cells, cell debris and nuclei are removed by centrifugation at $20,000 \times g$ for 20 minutes, and microsomal membranes are collected by centrifugation at $200,000 \times g$ for 1 hour. The membrane pellet is homogenized in storage buffer (25mM Bis-Tris-Propane, pH 7.0, 0.25M NaCl, 10% (w/v) glycerol and 1mM EDTA). Wax synthase activity is detected in these membranes using assay conditions as described for the jojoba enzyme. radiolabelled substrate is the same as for the jojoba 20 example (i.e. [1-14C] palmitoyl-CoA), however, 16:0 rather than 18:1 is used as the alcohol acceptor, and Bis-Tris-Propane buffer at pH 7.0 is utilized.

The Euglena wax synthase activity is solubilized by
incubation of the membranes with 2% CHAPS in the presence
of 0.5M NaCl. Solubilization of the protein is
demonstrated by the detection of enzyme activity in the
supernatant fraction after centrifugation at 200,000 x g
for 1 hour. The activity of the solubilized enzyme is
detected by dilution of the CHAPS concentration to ~0.3%
(i.e. to below its CMC). It is not necessary to
incorporate the enzyme into phospholipid vesicles as was
the case for the solubilized jojoba wax synthase.

For partial purification, the solubilized *Euglena* wax synthase activity is subjected to chromatographic separation on Blue A agarose medium. The column is equilibrated with 0.1M NaCl in a column buffer containing; 25mM Bis-Tris-Propane, pH 7.0, 20% (w/v) glycerol, 0.75% CHAPS and 1mM EDTA. The sample containing solubilized wax

synthase activity is diluted to 0.1M NaCl and loaded onto a 1 x 7cm column (5.5ml bed volume). The column is washed with equilibration buffer and subjected to a linear NaCl gradient (0.1M to 1.0M NaCl) in column buffer. Wax synthase activity is eluted as a broad peak in the last half of the salt gradient.

SDS-PAGE analysis of column fractions reveals that the polypeptide complexity of the activity eluted from the column is greatly reduced relative to the loaded material. A polypeptide with an apparent molecular mass of ~41kD was observed to track with wax synthase activity in the column fractions. Further purification techniques, such as described for jojoba and *Acinetobacter* are conducted to verify the association of wax synthase activity with the ~41kD peptide.

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For further analysis of wax synthase activity in Euglena, size exclusion chromatography was conducted as follows. A microsomal membrane preparation was obtained from Euglena cells grown on liquid, heterotrophic, medium 20 (Tani et al., supra) in the dark. Wax synthase activity was solubilized by treating the membranes with 2% (w/v)CHAPS and 500mM NaCl in a buffered solution (25mM Bis-Tris, pH 7.0, 1mM EDTA and 10% (w/v) glycerol) for 1 hour on ice. After dilution of the CHAPS to 0.75% and the NaCl to 200mM 25 by addition of a dilution buffer, the sample was centrifuged at $\sim 200,000 \times g$ for 1.5 hours. The supernatant fraction was loaded onto a Blue A dye column preequilibrated with Column Buffer (25mM Bis-Tris pH 7.0, 1mM EDTA, 10% glycerol, 0.75% CHAPS) which also contained 200mM NaCl. The column was washed with Column Buffer containing 200mM NaCl until the A280 of the effluent returned to the preload value. Wax synthase activity which had bound to the column was released by increasing the NaCl concentration in the Column Buffer to 1.5M. The fractions from the Blue A column containing wax synthase activity released by the 1.5M NaCl (~20ml combined volume) were pooled and concentrated approximately 30-fold via ultrafiltration (Amicon pressure cell fitted with a YM 30 membrane). The concentrated material from the Blue A

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column was used as the sample for a separation via size exclusion chromatography on Superose 12 medium (Pharmacia).

Approximately $200\mu l$ of the sample was loaded onto a Superose 12 column (HR 10/30), pre-equilibrated with Column Buffer containing 0.5M NaCl, and developed at a flow rate of 0.1ml/min. The wax synthase activity eluted from the column as a smooth peak. Comparison of the elution volume of the wax synthase activity with the elution profiles of molecular mass standard proteins yielded an estimate of

- 166kD for the apparent molecular mass of the enzyme. 10 Fractions which contained wax synthase activity were analyzed via SDS-polyacrylamide gel electrophoresis followed by silver staining. A preliminary analysis of the polypeptide profiles of the various fractions did not
- reveal any proteins with molecular masses of 100kD or 15 greater whose staining intensity appeared to match the activity profile. The wax synthase polypeptide may be present as a minor component in the sample mixture that is not readily detectable on the silver-stained gel.
- Alternatively, the enzyme may be composed of subunits which 20 are dissociated during SDS-PAGE.
- In addition to jojoba reductase, such as that encoded by the sequence provided in Figure 1, reductase proteins from other sources are also desirable for use in 25 conjunction with the wax synthase proteins of this Such proteins may be identified and obtained from organisms known to produce wax esters from alcohol and acyl substrates.
- 30 For example, an NADH-dependent fatty acyl-CoA reductase activity can be obtained from microsomal membranes isolated from Euglena gracilis. Methods which may be used to isolate microsomal membranes are described, for example in the published PCT patent application WO 92/14816 (application number PCT/US92/03164, filed February 35 21, 1992). The reductase activity is solubilized from these membranes using the same approaches as used for jojoba reductase and wax synthase. Membranes are incubated on ice for one hour with various amounts of the detergent,

CHAPS, in a buffering solution consisting of 25mM BisTris, pH 6.9, 250mM NaCl, 10% glycerol and 1 mM EDTA. is then centrifuged at $200,000 \times g$ for one hour, and the supernatant and pellet fractions assayed for NADH-dependent reductase activity using radiolabeled palmitoyl-CoA and NADH as substrates. A convenient assay for reductase activity is described in PCT patent application WO 92/14816. Incubation of the membranes with 0.3, 0.5 or 0.7 the supernatant fractions, indicative of solubilization of 10 the enzyme. If CHAPS is omitted during the incubation and centrifugation, all of the reductase activity is found in the pellet fraction. All of the samples are diluted tenfold in this same buffer solution prior to assaying in order to dilute the CHAPS present during the incubation. 15 The presence of CHAPS in the assay at levels above the CMC (approximately 0.5%(w/v) results in inhibition of enzyme activity. Stability of the reductase activity in up to 2% CHAPS may be improved by increasing the glycerol concentration in the buffering solution to 20%. Reductase 20 activity is recovered by dilution of the CHAPS to below the CMC.

25 Example 7 - Isolation of Nucleic Acid Sequences Isolation of nucleic acid sequences from cDNA libraries or from genomic DNA is described.

A. Construction of Jojoba cDNA Libraries

RNA is isolated from jojoba embryos collected at 80-90 days post-anthesis using a polyribosome isolation method, initially described by Jackson and Larkins (Plant Physiol. (1976) 57:5-10), as modified by Goldberg et al. (Developmental Biol. (1981) 83:201-217). In this procedure all steps, unless specifically stated, are carried out at 4°C. 10gm of tissue are ground in liquid nitrogen in a Waring blender until the tissue becomes a fine powder. After the liquid nitrogen has evaporated, 170ml of extraction buffer (200mM Tris pH 9.0, 160mM KC1, 25mM EGTA, 70mM MgC12, 1% Triton X-100, 05% sodium deoxycholate, 1mM

spermidine, 10mM ß-mercaptoethanol, and 500mM sucrose) is added and the tissue is homogenized for about 2 minutes. The homogenate is filtered through sterile miracloth and centrifuged at $12,000 \times g$ for 20 minutes. The supernatant is decanted into a 500ml sterile flask, and 1/19 volume of a 20% detergent solution (20% Brij 35, 20% Tween 40, 20% Noidet p-40 w/v) is added at room temperature. solution is stirred at 4°C for 30 minutes at a moderate speed and the supernatant is then centrifuged at 12,000 imes g for 30 minutes.

About 30ml of supernatant is aliquoted into sterile Ti 60 centrifuge tubes and underlaid with 7ml of a solution containing 40mM Tris pH 9.0, 5mM EGTA, 200mM KC1, 30mM MgC12, 1.8M sucrose, 5mM ß-mercaptoethanol. The tubes are filled to the top with extraction buffer, and spun at 15 60,000 rpm for 4 hours at 4°C in a Ti60 rotor. Following centrifugation, the supernatant is aspirated off and 0.5mlof resuspension buffer (40mM Tris pH 9.0, 5mM EGTA, 200mM KC1, 30mM MgCl₂, 5mM ß-mercaptoethanol) is added to each tube. The tubes are placed on ice for 10 minutes, after 20 which the pellets are thoroughly resuspended and pooled. The supernatant is then centrifuged at 120 \times g for 10 minutes to remove insoluble material. One volume of selfdigested 1mg/ml proteinase K in 20mM Tris pH 7.6, 200mM 25 EDTA, 2% N-lauryl-sarcosinate is added to the supernatant and the mixture incubated at room temperature for 30 minutes.

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RNA is precipitated by adding 1/10 volume of sodium acetate and 2 volumes of ethanol. After several hours at -20°C RNA is pelleted by centrifugation at 12,000 \times g at 4°C for 30 minutes. The pellet is resuspended in 10ml of TE buffer (10mM Tris, 1mM EDTA) and extracted with an equal volume of Tris pH 7.5 saturated phenol. The phases are separated by centrifuging at 10,000 x g for 20 minutes at 4°C . The aqueous phase is removed and the organic phase is re-extracted with one volume of TE buffer. The aqueous phases are then pooled and extracted with one volume of chloroform. The phases are again separated by

centrifugation and the aqueous phase ethanol precipitated as previously described, to yield the polyribosomal RNA.

Polysaccharide contaminants in the polyribosomal RNA preparation are removed by running the RNA over a cellulose column (Sigma-cell 50) in high salt buffer (0.5M NaCl, 20mM Tris pH 7.5, 1mM EDTA, 0.1% SDS). The contaminant binds to the column and the RNA is collected in the eluant. eluant fractions are pooled and the RNA is ethanol precipitated. The precipitated total RNA is then resuspended in a smaller volume and applied to an oligo d(T) cellulose column to isolate the polyadenylated RNA.

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Polyadenylated RNA is used to construct a cDNA library in the plasmid cloning vector pCGN1703, derived from the commercial cloning vector Bluescribe M13- (Stratagene Cloning Systems; San Diego, CA), and made as follows. polylinker of Bluescribe M13- is altered by digestion with BamHI, treatment with mung bean endonuclease, and blunt-end ligation to create a BamHI-deleted plasmid, pCGN1700. pCGN1700 is digested with EcoRI and SstI (adjacent restriction sites) and annealed with a synthetic linker having restriction sites for BamHI, PstI, XbaI, ApaI and Smal, a 5' overhang of AATT, and a 3' overhang of TCGA. The insertion of the linker into pCGN1700 eliminates the EcoRI site, recreates the SstI (also, sometimes referred to as "SacI" herein) site found in Bluescribe, and adds the new restriction sites contained on the linker. resulting plasmid pCGN1702, is digested with HindIII and blunt-ended with Klenow enzyme; the linear DNA is partially digested with PvuII and ligated with T4 DNA wax synthase in

plasmid cloning vector. Briefly, the cloning method for cDNA synthesis is as The plasmid cloning vector is digested with SstIand homopolymer T-tails are generated on the resulting 3'overhang stick-ends using terminal deoxynucleotidyl transferase. The tailed plasmid is separated from undigested or un-tailed plasmid by oligo(dA)-cellulose chromatography. The resultant vector serves as the primer

dilute solution. A transformant having the lac promoter

region deleted is selected (pCGN1703) and is used as the

for synthesis of cDNA first strands covalently attached to either end of the vector plasmid. The cDNA-mRNA-vector complexes are treated with terminal transferase in the presence of deoxyguanosine triphosphate, generating G-tails at the ends of the cDNA strands. The extra cDNA-mRNA complex, adjacent to the BamHI site, is removed by BamHI digestion, leaving a cDNA-mRNA-vector complex with a BamHI stick-end at one end and a G-tail at the other. complex is cyclized using an annealed synthetic cyclizing linker which has a 5' BamHI sticky-end, recognition 10 sequences for restriction enzymes NotI, EcoRI and SstI, and a 3' C-tail end. Following ligation and repair the circular complexes are transformed into $E.\ coli$ strain DH5lpha(BRL, Gaithersburg, MD) to generate the cDNA library. jojoba embryo cDNA bank contains between approximately 15 1.5×10^6 clones with an average cDNA insert size of approximately 500 base pairs.

Additionally, jojoba polyadenylated RNA is also used to construct a cDNA library in the cloning vector \$\frac{\text{2APII}/EcoRI}\$ (Stratagene, San Diego, CA). The library is constructed using protocols, DNA and bacterial strains as supplied by the manufacturer. Clones are packaged using Gigapack Gold packaging extracts (Stratagene), also according to manufacturer's recommendations. The cDNA library constructed in this manner contains approximately 1 x 106 clones with an average cDNA insert size of approximately 400 base pairs.

B. Polymerase Chain Reaction

Using amino acid sequence information, nucleic acid
sequences are obtained by polymerase chain reaction (PCR).
Synthetic oligonucleotides are synthesized which correspond
to the amino acid sequence of selected peptide fragments.
If the order of the fragments in the protein is known, such
as when one of the peptides is from the N-terminus or the
selected peptides are contained on one long peptide
fragment, only one oligonucleotide primer is needed for
each selected peptide. The oligonucleotide primer for the
more N-terminal peptide, forward primer, contains the
encoding sequence for the peptide. The oligonucleotide

primer for the more C-terminal peptide, reverse primer, is complementary to the encoding sequence for the selected peptide. Alternatively, when the order of the selected peptides is not known, two oligonucleotide primers are required for each peptide, one encoding the selected amino acid sequence and one complementary to the selected amino acid sequence. Any sequenced peptides may be selected for construction of oligonucleotides, although more desirable peptides are those which contain amino acids which are encoded by the least number of codons, such as methionine, tryptophan, cysteine, and other amino acids encoded by fewer than four codons. Thus, when the oligonucleotides are mixtures of all possible sequences for a selected peptide, the number of degenerate oligonucleotides may be low.

PCR is conducted with these oligonucleotide primers using techniques that are well known to those skilled in the art. Jojoba nucleic acid sequences, such as reverse transcribed cDNA, DNA isolated from the cDNA libraries described above or genomic DNA, are used as template in these reactions. In this manner, segments of DNA are produced. Similarly, segments of Acinetobacter w DNA are obtained from PCR reactions using oligonucleotide primers to the N-terminal and tryptic digest peptides described in Example 6A. The PCR products are analyzed by gel electrophoresis techniques to select those reactions yielding a desirable wax synthase fragment.

C. <u>Screening Libraries for Sequences</u>

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DNA fragments obtained by PCR are labeled and used as a probe to screen clones from the cDNA libraries described above. DNA library screening techniques are known to those in the art and described, for example in Maniatis et al. (Molecular Cloning: A Laboratory Manual, Second Edition (1989) Cold Spring Harbor Laboratory Press). In this manner, nucleic acid sequences are obtained which may be analyzed for nucleic acid sequence and used for expression of the plant cytoplasmic protein involved in fatty acyl-CoA metabolism in various hosts, both procaryotic and eucaryotic.

An approximately 1500 nucleotide jojoba cDNA clone is obtained in this manner. Comparison to the peptide fragments provided in Table 2 reveals the presence of each of these peptides in the translated sequence, with the exception of SQ1129. Northern analysis of jojoba embryo RNA indicates that the mRNA is approximately 2kb in length. Additional nucleic acid sequence is obtained using further PCR techniques, such as 5' RACE (Frohman et al., Proc. Nat. Acad. Sci. (1988) 85:8998-9002). Alternatively, additional sequences may be obtained by rescreening cDNA libraries or 10 from genomic DNA. Preliminary DNA sequence of a jojoba gene is presented in Figure 2. Further DNA sequence analysis of additional clones indicates that there are at least two classes of cDNA's encoding this jojoba protein. A plasmid containing the entire coding region in pCGN1703 15 is constructed to contain a SalI site approximately 8 nucleotides 5' to the ATG start codon, and is designated pCGN7614. The complete DNA sequence of pCGN7614 is presented in Figure 3. The major difference between the two classes of cDNAs as represented in the sequences in 20 Figures 2 and 3 is the presence (Figure 2) or absence (Figure 3) of the 6 nucleotide coding sequence for amino acids 23 and 24 of Figure 2.

25 D. Expression of Wax Synthase Activity in E. coli

The gene from pCGN7614 is placed under the control of the Tac promoter of $\it E.~coli$ expression vector pDR540 (Pharmacia) as follows. pCGN7614 DNA is digested at the SalI sites and the ends are partially filled in using the Klenow fragment of DNA polymerase I and the nucleotides TTP 30 and dCTP. The pDR540 vector is prepared by digesting with BamHI and partially filling in the ends with dGTP and dATP. The 1.8 kb fragment from pCGN7614 and the digested pDR540 vector are gel purified using low melting temperature agarose and ligated together using T4 DNA ligase. A colony 35 containing the encoding sequence in the sense orientation relative to the E. coli promoter was designated pCGN7620, and a colony containing the gene in the antisense orientation was designated pCGN7621.

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To assay for wax synthase activity, 50 ml cultures of pCGN7620 and pCGN7621 are grown to log phase in liquid culture, and induced for 2 hours by the addition of IPTG to a concentration of 1mM. The cells are harvested by centrifugation and subjected to the assay for wax synthase activity as described for jojoba extracts. TLC analysis indicates that the cell extract from pCGN7620 directs synthesis of wax ester, while the control extract from pCGN7621 does not direct the synthesis of wax ester. The wax synthase assay in these harvested cells was verified by a second assay, however, further attempts to produce wax synthase activity in E. coli cells transformed with reductase constructs have been unsuccessful.

15 Example 8 - Constructs for Plant Expression

Constructs which provide for expression of the plant cytoplasmic protein involved in fatty acyl-CoA metabolism and reductase sequences in plant cells may be prepared as follows.

20 A. <u>Expression Cassettes</u>

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Expression cassettes which contain 5' and 3' regulatory regions from genes expressed preferentially in seed tissues may be prepared from napin, Bce4 and ACP genes as described, for example in WO 92/03564.

For example, napin expression cassettes may be prepared as follows. A napin expression cassette, pCGN1808, which may be used for expression of wax synthase or reductase gene constructs is described in Kridl et al. (Seed Science Research (1991) 1:209-219), which is incorporated herein by reference.

Alternatively, pCGN1808 may be modified to contain flanking restriction sites to allow movement of only the expression sequences and not the antibiotic resistance marker to binary vectors such as pCGN1557 (McBride and Summerfelt, supra). Synthetic oligonucleotides containing KpnI, NotI and HindIII restriction sites are annealed and ligated at the unique HindIII site of pCGN1808, such that only one HindIII site is recovered. The resulting plasmid, pCGN3200 contains unique HindIII, NotI and KpnI restriction

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sites at the 3'-end of the napin 3'-regulatory sequences as confirmed by sequence analysis.

The majority of the napin expression cassette is subcloned from pCGN3200 by digestion with HindIII and SacI and ligation to HindIII and SacI digested pIC19R (Marsh, et al. (1984) Gene 32:481-485) to make pCGN3212. The extreme 5'-sequences of the napin promoter region are reconstructed by PCR using pCGN3200 as a template and two primers. flanking the SacI site and the junction of the napin 5'promoter and the pUC backbone of pCGN3200 from the pCGN1808 10 construct. The forward primer contains ClaI, HindIII, NotI, and KpnI restriction sites as well as nucleotides 408-423 of the napin 5'-sequence (from the EcoRV site) and the reverse primer contains the complement to napin 15 sequences 718-739 which include the unique SacI site in the 5'-promoter. The PCR was performed using a Perkin Elmer/Cetus thermocycler according to manufacturer's specifications. The PCR fragment is subcloned as a bluntended fragment into pUC8 (Vieira and Messing (1982) Gene 20 19:259-268) and digested with HincII to give pCGN3217. Sequence of pCGN3217 across the napin insert verifies that no improper nucleotides were introduced by PCR. The napin 5-sequences in pCGN3217 are ligated to the remainder of the napin expression cassette by digestion with ClaI and SacI 25 and ligation to pCGN3212 digested with ClaI and SacI. The resulting expression cassette pCGN3221, is digested with HindIII and the napin expression sequences are gel purified away and ligated to pIC20H (Marsh, supra) digested with HindIII. The final expression cassette is pCGN3223, which 30 contains in an ampicillin resistant background, essentially identical 1.725 napin 5' and 1.265 3' regulatory sequences as found in pCGN1808. The regulatory regions are flanked with HindIII, NotI and KpnI restriction sites and unique SalI, BglII, PstI, and XhoI cloning sites are located between the 5' and 3' noncoding regions. 35

Similarly, a cassette for cloning of sequences for transcription regulation under the control of 5' and 3' regions from an oleosin gene may be prepared. Sequence of a *Brassica napus* oleosin gene was reported by Lee and Huang

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(Plant Phys. (1991) 96:1395-1397). Primers to the published sequence are used in PCR reactions to obtain the 5' and 3' regulatory regions of an oleosin gene from Brassica napus cv. Westar. Two PCR reactions were performed, one to amplify approximately 950 nucleotides upstream of the ATG start codon for the oleosin gene, and one to PCR amplify approximately 600 bp including and downstream of the TAA stop codon for the oleosin gene. PCR products were cloned into plasmid vector pAMP1 (BRL) according to manufacturers protocols to yield plasmids 10 pCGN7629 which contains the oleosin 5' flanking region and pCGN7630 which contains the 3' flanking region. primers included convenient restriction sites for cloning the 5' and 3' flanking regions together into an expression 15 cassette. A PstI fragment containing the 5' flanking region from pCGN7629 was cloned into PstI digested pCGN7630 to yield plasmid pCGN7634. The BssHII (New England BioLabs) fragment from pCGN7634, which contains the entire oleosin expression cassette was cloned into BssHII digested 20 pBCSK+ (Stratagene) to provide the oleosin cassette in a plasmid, pCGN7636. Sequence of the oleosin cassette in pCGN7636 is provided in Figure 4. The oleosin cassette is flanked by BssHII, KpnI and XbaI restriction sites, and contains SalI, BamHI and PstI sites for insertion of wax synthase, reductase, or other DNA sequences of interest 25 between the 5' and 3' oleosin regions.

The gene sequences are inserted into such cassettes to provide expression constructs for plant transformation methods. For example, such constructs may be inserted into binary vectors for *Agrobacterium*-mediated transformation as described below.

B. <u>Constructs for Plant Transformation</u>

The plasmid pCGN7614 is digested with AflIII, and ligated with adapters to add BclI sites to the AflIII sticky ends, followed by digestion with SalI and BclI. The fragment containing the plant cytoplasmic protein involved in fatty acyl-CoA metabolism gene is gel purified and cloned into SalI/BamHI digested pCGN3223, a napin expression cassette. The resulting plasmid which contains

the plant cytoplasmic protein involved in fatty acyl-CoA metabolism gene in a sense orientation in the napin expression cassette is designated pCGN7624. DNA isolated from pCGN7624 is digested with Asp718 (a KpnI isoschizimer), and the napin/plant cytoplasmic protein involved in fatty acyl-CoA metabolism fusion gene is cloned into Asp718 digested binary vector pCGN1578 (McBride and Summerfelt, supra). The resultant binary vector, designated pCGN7626, is transformed into Agrobacterium strain EHA101 and used for transformation of Arabidopsis and rapeseed explants.

Additional binary vectors are prepared from pCGN1578, pCGN1559 and other vectors described by McBride et al. (supra) by substitution of the pCGN1578 and pCGN1559 linker regions with a linker region containing the following restriction digestion sites:

Asp718/AscI/PacI/XbaI/BamHI/SwaI/Sse8387 (PstI)/HindIII. This results in pCGN1578PASS or pCGN1559PASS, and other modified vectors which are designated similarly. AscI, PacI, SwaI and Sse8387 have 8-base restriction recognition sites. These enzymes are available from New England BioLabs: AscI, PacI; Boehringer Manheim: SwaI and Takara (Japan): Sse8387.

C. Reductase Constructs for Plant Transformation Constructs for expression of reductase in plant cells using 5' and 3' regulatory regions from a napin gene, are prepared.

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A reductase cDNA (in the pCGN1703 vector described above) designated pCGN7571, is digested with SphI (site in 3' untranslated sequence at bases 1594-1599) and a SalI linker is inserted at this site. The resulting plasmid is digested with BamHI and SalI and the fragment containing the reductase cDNA gel purified and cloned into BglII/XhoI digested pCGN3223, the napin cassette described above, resulting in pCGN7585.

A *HindIII* fragment of pCGN7585 containing the napin 5'/reductase/napin 3' construct is cloned into HindIII digested pCGN1578 (McBride and Summerfelt, *supra*), resulting in pCGN7586, a binary vector for plant transformation.

Plant transformation construct pCGN7589, also containing the jojoba reductase gene under expression of a napin promoter, is prepared as follows. pCGN7571 is in vitro mutagenized to introduce an NdeI site at the first ATG of the reductase coding sequence and a BglII site immediately upstream of the NdeI site. BamHI linkers are introduced into the SphI site downstream of the reductase coding region. The 1.5 kb BglII-BamHI fragment is gel purified and cloned into BglII-BamHI digested pCGN3686 (see below), resulting in pCGN7582.

pCGN3686 is a cloning vector derived from Bluescript KS+ (Stratagene Cloning Systems; San Diego, CA), but having a chloramphenicol resistance gene and a modified linker region. The source of the chloramphenicol resistance gene, pCGN565 is

a cloning vector based on pUC12-cm (K. Buckley Ph.D. Thesis, Regulation and expression of the phi X174 lysis gene, University of California, San Diego, 1985), but containing pUC18 linkers (Yanisch-Perron, et al., Gene (1985) 53:103-119). pCGN565 is digested with HhaI and the fragment

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- containing the chloramphenicol resistance gene is excised, blunted by use of mung bean nuclease, and inserted into the EcoRV site of Bluescript KS- (Stratagene: La Jolla, CA) to create pCGN2008. The chloramphenicol resistance gene of pCGN2008 is removed by EcoRI/HindIII digestion. After
- treatment with Klenow enzyme to blunt the ends, the fragment is ligated to DraI digested Bluescript KS+. A clone that has the DraI fragment containing ampicillin resistance replaced with the chloramphenical resistance is chosen and named pCGN2015. The linker region of pCGN2015 is modified to provide pCGN3686, which contains the following restriction digestion sites 51 to 31 in the laggetime.
 - digestion sites, 5' to 3' in the lacZ linker region: PstI, BglII, XhoI, HincII, SalI, HindIII, EcoRV, EcoRI, PstI, SmaI, BamHI, SpeI, XbaI and SacI.

An XhoI linker is inserted at the XbaI site of pCGN7582.

The BglII-XhoI fragment containing the reductase gene is isolated and cloned into BglII-XhoI digested pCGN3223. The resulting plasmid, which lacks the 5' untranslated leader sequence from the jojoba gene, is designated pCGN7802. The napin/reductase fragment from pCGN7802 is excised with

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 ${\it Hin}{\it dIII}$ and cloned into ${\it Hin}{\it dIII}$ digested pCGN1578 to yield pCGN7589.

An additional napin/reductase construct is prepared as follows. The reductase cDNA pCGN7571 (Figure 1) is mutagenized to insert SalI sites 5' to the ATG start codon (site is 8 base pairs 5' to ATG) and immediately 3' to the TAA translation stop codon, resulting in pCGN7631. pCGN7631 is digested with SalI and the approximately 1.5 kb fragment containing the reductase encoding sequence is cloned into SalI/XhoI digested napin cassette pCGN3223. A resulting plasmid containing the reductase sequence in the sense orientation is designated pCGN7640. pCGN7640 is digested with HindIII, and the fragment containing the oleosin/reductase construct is cloned into HindIII digested binary vector pCGN1559PASS, resulting in binary construct pCGN7642.

A construct for expression of reductase under control of oleosin regulatory regions is prepared as follows. The reductase encoding sequence is obtained by digestion of pCGN7631 with SalI, and ligated into SalI digested pCGN7636, the oleosin cassette. A resulting plasmid containing the reductase sequence in the sense orientation is designated pCGN7641. pCGN7641 is digested with XbaI, and the fragment containing the oleosin/reductase construct is cloned into XbaI digested binary vector pCGN1559PASS, resulting in binary construct pCGN7643.

Binary vector constructs are transformed into Agrobacterium cells, such as of strain EHA101 (Hood et al., J. Bacteriol (1986) 168:1291-1301), by the method of Holsters et al. (Mol. Gen. Genet. (1978) 163:181-187) and used in plant transformation methods as described below.

Example 9 - Plant Transformation Methods

A variety of methods have been developed to insert a

35 DNA sequence of interest into the genome of a plant host to
obtain the transcription or transcription and translation
of the sequence to effect phenotypic changes.

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Brassica Transformation

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Seeds of high erucic acid, such as cultivar Reston, or Canola-type varieties of Brassica napus are soaked in 95% ethanol for 2 min. surface sterilized in a 1.0% solution of sodium hypochlorite containing a drop of Tween 20 for 45 min., and rinsed three times in sterile, distilled water. Seeds are then plated in Magenta boxes with 1/10th concentration of Murashige minimal organics medium (Gibco; Grand Island, NY) supplemented with pyriodoxine (50 μ g/1), nicotinic acid (50 μ g/1), glycine (200 μ g/1), and 0.6% Phytagar (Gibco) pH 5.8. Seeds are germinated in a Percival chamber at 22°C. in a 16 h photoperiod with cool fluorescent and red light of intensity approximately 65 μ Einsteins per square meter per second (μ Em-2s-1).

Hypocotyls are excised from 5-7 day old seedlings, cut 15 into pieces approximately 4mm in length, and plated on feeder plates (Horsch et al., Science (1985) 227:1229-1231). Feeder plates are prepared one day before use by plating 1.0ml of a tobacco suspension culture onto a petri plate ($100 \times 25 mm$) containing about 30ml MS salt base (Carolina Biological, Burlington, NC) 100mg/l inositol, 1.3mg/l thiamine-HCl, 200mg KH_2PO_4 with 3% sucrose, 2,4-D (1.0mg/l), 0.6% w/v Phytagar, and pH adjusted to 5.8 prior to autoclaving (MS 0/1/0 medium). A sterile filter paper disc (Whatman 3mm) is placed on top of the feeder layer 25 prior to use. Tobacco suspension cultures are subcultured weekly by transfer of 10ml of culture into 100ml fresh MS medium as described for the feeder plates with 2,4-D (0.2mg/1), Kinetin (0.1mg/1). In experiments where feeder cells are not used hypocotyl explants are cut and placed 30 onto a filter paper disc on top of MSO/1/0 medium. All hypocotyl explants are preincubated on feeder plates for 24 h. at 22°C in continuous light of intensity $30\mu\text{Em}^{-2}\text{S}^{-1}$ to $65\mu EM^{-2}S^{-1}$.

Single colonies of A. tumefaciens strain EHA101 containing a binary plasmid with the desired gene construct are transferred to 5ml MG/L broth and grown overnight at 30°C. Hypocotyl explants are immersed in 7-12ml MG/L broth with bacteria diluted to 1x10⁸ bacteria/ml and after 10-25

min. are placed onto feeder plates. Per liter MG/L broth contains 5g mannitol, 1g L-Glutamic acid or 1.15g sodium glutamate, 0.25g kH₂PO₄, 0.10g NaCl, 0.10g MGSO₄·7H₂O, 1mg biotin, 5g tryptone, and 2.5g yeast extract, and the broth is adjusted to pH 7.0. After 48 hours of co-incubation with Agrobacterium, the hypocotyl explants are transferred to B5 0/1/0 callus induction medium which contains filter sterilized carbenicillin (500mg/l, added after autoclaving) and kanamycin sulfate (Boehringer Mannheim; Indianapolis, IN) at concentrations of 25mg/l.

After 3-7 days in culture at 65µEM-2S-1 continuous light, callus tissue is visible on the cut surface and the hypocotyl explants are transferred to shoot induction medium, B5BZ (B5 salts and vitamins supplemented with 3mg/l benzylaminopurine, 1mg/l zeatin, 1% sucrose, 0.6% Phytagar and pH adjusted to 5.8). This medium also contains carbenicillin (500mg/l) and kanamycin sulfate (25mg/l). Hypocotyl explants are subcultured onto fresh shoot induction medium every two weeks.

Shoots regenerate from the hypocotyl calli after one to three months. Green shoots at least 1cm tall are excised from the calli and placed on medium containing B5 salts and vitamins, 1% sucrose, carbenicillin (300mg/l), kanamycin sulfate (50mg/l) and 0.6% w/v Phytagar). After 2-4 weeks shoots which remain green are cut at the base and transferred to Magenta boxes containing root induction medium (B5 salts and vitamins, 1% sucrose, 2mg/l indolebutyric acid, 50mg/l kanamycin sulfate and 0.6% Phytagar). Green rooted shoots are tested for thioesterase activity.

Arabidposis Transformation

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Transgenic Arabidopsis thaliana plants may be obtained by Agrobacterium-mediated transformation as described by Valverkens et al., (Proc. Nat. Acad. Sci. (1988) 85:5536-5540). Constructs are transformed into Agrobacterium cells, such as of strain EHA101 (Hood et al., J. Bacteriol (1986) 168:1291-1301), by the method of Holsters et al. (Mol. Gen. Genet. (1978) 163:181-187).

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Peanut Transformation

DNA sequences of interest may be introduced as expression cassettes, comprising at least a promoter region, a gene of interest, and a termination region, into a plant genome via particle bombardment.

Briefly, tungsten or gold particles of a size ranging from 0.5mM-3mM are coated with DNA of an expression cassette. This DNA may be in the form of an aqueous mixture or a dry DNA/particle precipitate.

Tissue used as the target for bombardment may be from cotyledonary explants, shoot meristems, immature leaflets, or anthers. The bombardment of the tissue with the DNA-coated particles is carried out using a Biolistics particle gun (Dupont; Wilmington, DE). The particles are placed in the barrel at variable distances ranging from 1cm-14cm from the barrel mouth. The tissue to be bombarded is placed beneath the stopping plate; testing is performed on the tissue at distances up to 20cm. At the moment of discharge, the tissue is protected by a nylon net or a combination of nylon nets with mesh ranging from 10mM to 300mM.

Following bombardment, plants may be regenerated following the method of Atreya, et al., (Plant Science Letters (1984) 34:379-383). Briefly, embryo axis tissue or cotyledon segments are placed on MS medium (Murashige and Skoog, Physio. Plant. (1962) 15:473) (MS plus 2.0 mg/l 6benzyladenine (BA) for the cotyledon segments) and incubated in the dark for 1 week at 25 \pm 2°C and are subsequently transferred to continuous cool white fluorescent light (6.8 W/m^2). On the 10th day of culture, the plantlets are transferred to pots containing sterile soil, are kept in the shade for 3-5 days are and finally moved to greenhouse. The putative transgenic shoots are Integration of exogenous DNA into the plant genome may be confirmed by various methods know to those skilled in the art.

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Example 10 - Analysis of Transformed Plants for Wax Production

Seeds or other plant material from transformed plants may be analyzed for wax synthase activity using the wax synthase assay methods described in Example 1.

Plants which have both the reductase and wax synthase constructs are also assayed to measure wax production. Such plants may be prepared by Agrobacterium transformation methods as described above. Plants having both of the desired gene constructs may be prepared by cotransformation with reductase and wax synthase constructs or by combining the wax synthase and reductase constructs on a single plant transformation binary vector. In addition, re-transformation of either wax synthase expressing plants or reductase expressing plants with constructs encoding the other desired gene sequence may also be used to provide such reductase and wax synthase expressing plants. Alternatively, transgenic plants expressing reductase produced by methods described herein may be crossed with plants expressing wax synthase which have been similarly produced. In this manner, known methods of plant breeding are used to provide reductase and wax synthase expressing transgenic plants.

Such plants may be assayed for the presence of wax esters, for example by separation of TAG from wax esters as described by Tani et al. (supra). GC analysis methods may be used to further analyze the resulting waxes, for example as described by Pina et al. (Lipids (1987) 22(5):358-361.

The above results demonstrate the ability to obtain partially purified wax synthase proteins which are active in the formation of wax esters from fatty alcohol and fatty acyl substrates. Methods to obtain the wax synthase proteins and amino acid sequences thereof are provided. In addition wax synthase nucleic acid sequences obtained from the amino acid sequences are also provided. These nucleic acid sequences may be manipulated to provide for transcription of the sequences and/or expression of wax synthase proteins in host cells, which proteins may be used

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for a variety of applications. Such applications include the production of wax ester compounds when the wax synthase is used in host cells having a source of fatty alcohol substrates, which substrates may be native to the host cells or supplied by use of recombinant constructs encoding a fatty acyl reductase protein which is active in the formation of alcohols from fatty acyl substrates.

Example 11 - Analysis of Transformed Plants for 10 VLCFA Production

Seeds from transformed plants are analyzed by gas chromatography (GC) for fatty acid content. The following tables provide breakdowns of fatty acids on a percentage basis, demonstrating altered VLCFA production in plants transformed with binary vector pCGN7626 (Example 8).

Table 3

percentage of fatty acids for each plant and fatty some transformed by pCGN7626, showing saturation. Twenty seeds were pooled Seeds from canola plants, some transform of a given carbon chain length:saturation. acids determined by gas chromatography. Control canola plants (plants 1 and 2) of Table 3 contain less than 2% VLCFA in their seed oil. Plants 3 through 20 in Table 3 are transgenic. The majority (14/18) of the plants transformed with pCGN7626 have significantly higher levels of VLCFA. The VLCFA for the highly transformed with pCGN7626 have significantly higher levels of VLCFA. The VLCFA for expresssing transgenics range from about 5% to about 22% of the total fatty acids.

:2	0.00	0.00	딩	0.66	21	02	17	8	02	8	17	02	00	딩	02	8	ĺŠ	Ħ	22	12
\$ 22:	0	0	0.0	0.	0.23	0.02	0.0	0.00	0.02	0.08	0.0	0.02	0.00	0.0	0.0	0.08	0.0	0.0	0.02	0.02
\$ 22:1	0.01	0.01	0.47	4.84	1.73	0.88	0.49	0.01	1.39	0.88	0.00	0.46	0.01	0.00	0.46	0.69	0.25	0.44	0.26	0.58
\$ 22:0	0.24	0.25	0.24	0.39	0.31	0.27	0.34	0.31	0.24	0.24	0.25	0.28	0.26	0.34	0.26	0.33	0.26	0.26	0.22	0.15
\$ 20:2	0.08	0.09	0.33	1.11	0.67	0.47	0.35	0.09	0.53	0.48	0.10	0.41	0.11	0.04	0.34	0.47	0.24	0.31	0.25	0.41
\$ 20:1	1.20	1.31	4.97	14.27	9.75	6.93	5.41	1.27	7.24	6.72	1.25	4.88	1.35	1.17	4.19	5.03	3.86	5.13	3.77	4.48
\$ 20:0	0.45	0.41	0.46	0.49	0.49	0.46	0.44	0.45	0.45	0.44	0.41	0.39	0.43	0.39	0.39	0.47	0.47	0.43	0.43	0.36
\$ 18:3	12.48	11.25	15.95	14.57	14.89	13.74	14.90	11.20	16.15	15.52	16.83	17.50	14.35	15.39	19.78	15.51	14.89	15.20	15.09	22.87
\$ 18:2	21.14	22.09	19.24	19.60	18.76	20.34	19.40	19.52	20.51	20.48	21.44	22.28	21.08	20.93	20.65	23.86	20.04	19.57	19.77	20.15
\$ 18:1	58.42	58.89	52.01	38.12	46.74	51.00	52.36	60.63	47.57	48.91	53.17	48.04	56.23	53.08	47.06	46.98	53.62	52.20	53.74	44.57
\$ 18:0	1.30	1.12	1.11	0.76	06.0	0.95	0.99	1.10	0.91	0.93	1.16	0.94	1.07	0.88	0.89	0.93	1.26	1.02	1.14	0.92
ON	1	2	3	4	5	9	7	8	6	10	11	12	13	14	15	16	17	18	19	20

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Table 4

Canola plants, some transformed by pCGN7626, showing percentage of fatty acids of a given carbon chain length:saturation.

transformed by pCGN7626, showing inheritance of the altered VLCFA phenotype. One plant, plant 11, did not inherit the altered phenotype. This plant also did not show inheritance of the Plants 1 and 2 in Table 4 are controls. Plant 3 is a repeat of plant 4 of Table 3. Plants 4 through 13 are seed of plants grown out from the seed of a single canola plant transgene by a Kan germination assay.

ſ		1-	ना	51	7	TE	नार	61	~	77	31	=	Τ~	<u> </u>	<u> </u>	. Т.	_	1-
	824:1	6		7.0	0.67	0		0.30	0.4	2,0		0.24	0	20.00		00.00	0.59	0 37
	824:0	0	7 6	0.01	0.24	0	2 6	777	0.04	5	70.0	0.21	0	100		20.0	0.70	0.04
	\$22:2	0		3	0.58	0.22	20	0.43	0.20	0 17	;	0.09	0.01	12	200	200	0.23	0.32
	\$22:1	0	0.00		3.93	1.78	1 76	7	1.56	1 27		0.84	0.27	1	2 6	7 2 2	7 . / 0	1.83
000	\$77:O	0.25	0 26	2 6	0.39	0.34	2	1	0.29	0.29		0.31	0.24	0.33	α 0	200	0.47	0.34
000	2:079	0.08	0	3	1.05	0.63	0.76		0.00	0.64	,	0.47	0.24	0.54	0	6	30.0	0.79
9.20.1	T:079	1.19	1.30	2 2 2	12.31	7.70	8.83	,	8.01	7.80	60	0.03	3.48	7.68	1.18	7 5.8	?	7.62
0.000	920:0	0.43	0.42	1	0.31	0.50	0.46	7	0.45	0.46	0	0.33	0.39	0.55	0.41	0 50		0.47
210.3	C:070	11.87	10.71	15.00					12.21	16.31	11 36	7.70	13.22	13.53	14.91	14.04		14.92
910.0	2.010	21.61	22.38	70 37	40.3	20.97	23.36		67.77	22.15	70 21	20:03	23.14	21.21	24.05	23.03		24.20
\$18.1	ı	58.14	58.73	۲	-	43.21	42.48	00 77		43.13	18 73		52.27	46.79	51.73	44.56	100	41.32
818.0	2	1.25	1.02	C	20:	0.98	0.87	0 07		0.96	1 17	1	0.97	1.12	0.98	1.10	30	0.88
CZ		T	2	~	,	4	S	v	,	7	α	,	5	10	11	12	;	7

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The results of measurements of seeds of HEAR plants, controls and pCGN7626 transgenic, Pools of twenty seeds were analyzed by GC. evaluated for VLCFA content.

comprising about 0.1 to 0.5%. The results show significant alteration of the VLCFA patterns. HEAR (variety Reston) has 22:1 levels between 33 and 41 percent of its fatty acids with 24:1 Control Plants 3, 4, 7, 12-14 and 16-19 particularly showed an increase in 24:1 content, with one The remaining plants are transgenic. transgenic plant showing a 24:1 level of 2.7% of the seed oil plants. Plants 1 and 2 are control HEAR

\$24		,			2.69	1.21	0		7.7.	1.41	0.09	0.13	0	7.0	,	1.43	1.39	1.58	0		?;	1.46	1.30	1 85
\$24:0		000	3 5	70.0	0.06	0.05	0	0.05	200	20.0	00.0	0.04	0.02	200	3 5	20.0	0.02	0.05	0	20.0	0.00	0.02	0.04	0.03
\$22:2		0 7 0		0.40	1.72	1.16	1.22	1 27	10	20.00	0.62	0.96	0.59	0 73	200	25.0	0.72	1.16	0.02	1 56	000	77.7	1.10	1.24
\$22:1		40 57			38.32	37.84	37.16	38 29		27.50	37.02	36.48	34.55	35 82	26.26	200	22.3	35.69	0.78	30 10	36.25	0 20		38.53
\$22:0		0.48	a C	200		0.54	0.53	0.47	0 44			0.61	90.0	0.37	0 47			0.43	0.17	0 77	5		0.0	0.68
\$20:2		0.75	c		اد	0.90	0.95	0.93			0 0	0.7	0.72	0.84	0	0		/8.0	0.54	0.78	70	200	0.0	0.83
\$20:1		6.00	8.36	200	3.44	6.60	6.32	6.49	6.68	7 51	100	0.00	8.48	5.85	7.23	6 97			5.88	6.30	6 10	7 17		(1.15
\$20:0		0.46	0.46	70	7.7	0.48	0.42	0.44	0.48	0.44	2 2	000	0.51	0.35	0.46	0.47		7	0.35	0.45	0.51			0.53
\$18:3		12.32	9.74	1		• 1	12.77	11.26	11.73	10.60		60.17	10.25	12.52	10.10	10.01	10 07	10:26	16.95	10.86	10.79	0 42		11.43
\$18:2		18.07	18.49	17 45	71.05	• •	19.55	19.29	18.35	18.67			18.22	20.64	18.19	19.65	18 67	100	77.48	16.48	19.23	18.31		10.30
\$18:1		13.69	19.90	12 94	1	•	13.85	14.56	15.03	16.14	17 00	200	2 2	14.36	17.10	17.99	16 02		3	14.92	15.40	16.35		70.51
\$18:0	- 4	0.90	1.03	1.06	900		1.05	1.04	1.03	1.02			70.7	0.92	0.99	0.95	0.87	5	70.7	0.94	0.93	1.04	00	
<u>8</u>		7	2	3	-	r L		٥	7	8	6	, ,	3	T	12	13	14	100		19	17	18	0	7

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Table 6

Arabidopsis thaliana typically has The oil composition of plants transformed with pCGN7626 (plants 4-12) is shifted towards the longer chain fatty acids at the expense of 20:1. The 20:1 in transgenic plants decreased to as low as 15.5% while the 22:1 percentage increased to as high as 7.5%. In one transgenic seed oil with 21% 20:1 fatty acid, 2% 22:1 fatty acid, 0.02% 24:1 fatty acid (control plants plant the 24:1 content increased to 1.6% of the total fatty acids in the seed oil. Arabidopsis thaliana plants transformed with pCGN7626.

In Table 7 oil seed analysis results are given for T3 Brassica plants, (LEAR variety 212) transformed with pCGN7626

ON.	%18:0	818:1	\$18:2	%18:3	\$20:0	\$20:1	\$20:2	\$22:0	822:1	\$22:2	\$24:0	\$24.1
1			26.82	18.08	2.17		2.03	0.33	2.07	0.04	0.01	0
2				18.61	2.22	l .	1.83	0.26	1.80	0.02	0.01	0
m				18.30	2.07		2.02	0.10	2.00	0.02	0.05	0.0
4	1	ł		18.67	1.99		1.77	90.0	1.58	0.02	0.05	0.03
2		- 1		20.80	1.85		1.97	0.85	4.03	0.32	0.07	0.74
9	- 1			20.19	1.97	1	1.97	0.74	3.36	0.04	0.51	0.42
7				18.80	1.84		1.64	0.04	1.92	0.01	0.02	0.04
8				20.56	1.56		1.80	1.29	5.72	0.69	1-1	1 55
6	3.34	15.11	25.89	19.48	2.05	19.58	2.03	0.44	2.60	0.12	0.03	0.04
10				20.51	1.83		2.01	06.0	3.98	0.40	0.84	0 67
11				18.45	1.55		1.84	1.49	7.47	0.73	60 0	104
12	1.94			19.91	1.42	,	1.44	1.34	6.40	0.43	1.06	1 60

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NO I	STRAIN ID	\$16:0	\$16:1	\$18:0	\$18:1	\$18:2	*18:3	\$20:0	\$20:1	\$20:2	\$22:0	\$22:1	\$22:2	\$24:0	\$24:1	>18
` ↔	RESTON	2.54	0.05	0.79	17.54	12.12	9.59	0.54	8.80	0.49	0.55	46.13	0.38	0.00	0.08	56.97
7	RESTON	2.68	0.12	0.78	19.96	11.79	8.80	0.52	9.98	0.45	0.46	42.84	0.05	0.03	0.92	55.25
m	RESTON	2.59	0.12	0.73	19.15	11.96	7.90	0.46	8.40	0.41	0.38	47.30	90.0	0.00	0.10	57.11
4	RESTON	2.49	0.09	0.83	16.37	11.98	10.22	0.50	8.49	0.52	0.52	46.23	0.48	0.06	0.86	57.66
Ŋ	RESTON	2.65	0.15	0.81	17.63	14.18	6.51	0.43	7.80	0.35	0.40	46.87	0.46	0.00	1.21	57.52
9	RESTON	2.52	0.10	0.79	17.50	11.61	10.35	0.49	8.50	0.52	0.67	45.07	0.34	0.12	1.02	56.73
7	RESTON	2.84	0.20	0.73	17.86	11.60	9.18	0.44	9.51	0.46	0.30	45.97	0.21	0.00	0.18	57.07
œ	RESTON	2.71	0.14	0.81	17.64	12.09	11.15	0.50	8.56	0.54	09.0	43.46	0.39	0.10	0.81	54.96
6	RESTON	2.46	0.10	0.84	22.84	9.72	6.50	0.56	9.30	0.31	0.50	45.02	0.20	0.00	1.15	57.04
10	10 RESTON	2.57	0.13	0.78	23.40	9.80	6.41	0.53	8.83	0.36	0.38	45.28	0.15	0.00	0.86	56.39
11	11 7626-212-2-1	-1 2.92	0.15	0.64	25.92	10.42	6.85	0.46	15.21	0.61	0.92	28.79	1.33	0.45	7.78	55.55
12	12 7626-212-2-1	-1 3.05	0.28	0.74	29.57	11.37	6.94	0.56	17.72	0.65	77.0	22.67	0.77	0.11	4.43	47.68
13	13 7626-212-2-1	.1 2.80	0.12	0.52	19.06	11.56	8.73	0.41	13.78	0.77	0.67	33.64	1.45	0.00	5.44	56.16
14	14 7626-212-2-1	.1 2.88	0.25	0.76	20.92	11.12	5.38	0.58	11.50	0.48	1.19	34.51	1.26	0.65	7.79	57.96
15	15 7626-212-2-1	1 3.14	0.23	0.99	26.29	11.02	8.18	0.65	19.12	0.76	0.82	24.17	1.07	0.00	3.06	49.65
16	16 7626-212-2-1	1 2.83	0.23	0.77	28.54	10.55	7.50	0.67	18.72	0.62	0.93	23.40	0.98	0.31	3.48	49.11
17	17 7626-212-2-	1 2.82	0.15	0.68	23.05	10.65	6.93	0.53	16.81	0.70	0.88	28.46	1.25	0.08	6.41	55.12
18	18 7626-212-2-1	1 2.59	0.17	0.69	22.36	11.75	9.63	0.56	15.58	0.82	0.97	29.52	1.26	0.19	3.48	52.38
19	7626-212-2-1	1 2.46	0.15	0.71	21.51	11.35	9.03	0.54	13.52	0.64	0.78	33.54	1.14	0.15	3.87	54.18
		1 3.07	0.18	0.69	28.80	13.12	9.24	0.40	17.80	0.78	0.45	20.33	0.88	0.00	3.39	44.03
		2 3.36	0.30	0.83	25.51	14.30	10.62	0.44	14.30	0.75	0.39	26.58	0.61	0.00	1.48	44.55
			0.15	0.92	25.00	12.47	8.23	0.59	16.69	0.69	0.43	28.65	0.59	0.01	1.82	49.47
23	7626-212-2-2	2 2.62	0.11	98.0	21.14	12.45	11.23	0.54	16.50	06.0	0.48	29.92	98.0	0.07	1.72	50.99
24	7626-212-2-2		0.24	0.81	24.25	12.09	10.84	0.53	15.83	0.76	0.38	27.79 (99.0	0.07	1.99	48.01
52	7626-212-2-2	3.44	0.13	1.12	35.66	14.49	10.23	0.61	16.32	0.59	0.46	14.47 (0.14	0.05	1.67	34.31

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8 ¦	NO STRAIN ID	\$16:0	\$16:1	1 \$18:0	3 %18:1	\$18:2	\$18:3	\$20:0	\$20:1	\$20:2	\$22:0	\$22:1	\$22:2	\$24:0	\$24:1	>18
26	26 7626-212-2-2	2 2.90	0.22	0.79	20.44	13.05	11.06	0.43	12.54	0.68	00.0	35.58	0.09	0.02	1 60	50 07
27	27 7626-212-2-2	2 2.59	0.08	0.69	16.89	11.94	9.99	0.50	10.67	0.77	0.77	39.93	1.40	0.14	3 22	57.72
28	7626-212-2-2	2 2.80	0.12	0.82	21.71	12.94	9.73	0.61	14.96	0.90	0.72	30.39	1.04	0.00	22.0	2
29	7626-212-2-2	3.41	0.15	1.07	36.19	15.14	10.55	0.46	17.10	0.57	0.08	14.66	0.00	0.00	0.10	32 07
30	7626-212-2-2	2.97	0.11	0.96	24.24	13.21	9.58	0.58	15.50	0.84	0.56	26.59	3.09	0.00	1.60	48 76
31	7626-212-2-3		0.12	0.87	24.30	11.93	9.40	0.53	10.45	0.46	0.58	35.32	0.50	0.06	20.7	00.01
32			0.12	0.94	23.18	11.13	7.34	0.64	10.98	0.34	0.41	40.76	0.06	0.00	0.97	54 16
33	7626-212-2-3		0.18	2.28	23.96	11.50	8.17	0.49	8.80	0.53	0.57	36.37	0.41	0.07	1.96	49 20
34	7626-212-2-3	3.22	0.13	1.74	39.52	13.91	7.96	0.71	16.79	0.26	0.24	14.33	0.03	0.00	0.70	33.05
35	35 7626-212-2-3		0.00	1.74	26.41	11.98	4.23	1.15	11.37	0.47	0.84	36.39	0.08	0.00	1.68	51.00
36	7626-212-2-3	3.81	0.20	1.49	37.32	15.55	9.58	0.65	16.61	0.55	0.05	13.35	0.01	00.00	0.16	31 28
37	7626-212-2-3	2.88	0.16	1.37	25.49	12.95	8.90	0.69	14.10	0.58	0.35	30.54	0.11	0.02	1.25	47.64
38	7626-212-2-3	3.47	0.13	1.37	22.30	14.75	11.27	0.68	10.43	0.45	0.48	33.74	0.20	0.07	0.14	46.19
39	39 7626-212-2-3	3.61	0.18	1.98	29.46	11.76	5.03	1.17	13.56	0.36	0.74	29.88	0.18	0.00	1.42	47.33
40	40 7626-212-2-3	2.17	0.12	1.06	20.51	13.59	11.14	09.0	10.57	0.32	0.45	36.98	90.0	0.07	1 05	40.02
41	7626-212-2-4	2.71	0.15	0.74	16.79	14.51	10.60	0.51	9.40	0.89	0.67	37.72	1.22	0.06	75.4	53 63
42	7626-212-2-4	3.07	0.26	0.80	17.32	13.47	10.23		10.91	0.85		36.07	1.31	90.0	3.77	54.02
43	7626-212-2-4	3.00	0.09	0.94	23.10	15.70	9.32	0.52	16.33	0.92			0.73	0.07	2.62	47 19
44	44 7626-212-2-4	2.77	0.11	09.0	19.54	14.82	6.57	0.32	13.32	0.89			1.51	0.29	7 39	55 21
45	7626-212-2-4	2.87	0.14	96.0	17.40	14.75	9.39	99.0	7.58				0.72	0.10		53.83
46	7626-212-2-4	2.86	0.25	0.63	15.72	14.40	10.12	0.40	8.99				1.10	0.00		55.41
47	47 7626-212-2-4	3.30	0.18	96.0	18.64	14.78	14.88	0.36	13.37	0.76		31.24 (.18	0.00		45.99
48	7626-212-2-4	3.10	0.21	0.93	20.82	14.19	6.07	0.62	10.33			37.79	.70	0.09		54 46
49	7626-212-2-4	3.70	0.10	0.91	16.43	15.05	13.39	0.52	10.59				1.26			49.53
20	50 7626-212-2-4	3.10	0.24	1.69	29.12	12.66	6.21	1.06 1	14.43			_				46.35

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Analysis of T3 seed oil from LEAR plants transformed with the jojoba CE shows that up to 7.8 % of the seed oil is 24:1. As is seen from the controls, the Reston plants, which are HEAR, typically have only about 1% or less 24:1.

These data clearly show that the plant cytoplasmic protein involved in fatty acyl-CoA metabolism encoded by pCGN7626 can markedly alter the fatty acid composition of seed oil from several plant species. In plants that do not accumulate VLCFA, pCGN7626 causes the accumulation of significant quantities of VLCFA. In plants that do accumulate VLCFA, pCGN7626 shifts the fatty acid composition towards longer VLCFA.

When searching protein data bases for the jojoba protein sequence disclosed herein, a large region of homology was found between the jojoba encoded protein and 15 stilbene, reservatrol, and chalcone synthase. Stilbene, reservatrol and chalcone synthases are very similar to each other, catalyzing multiple condensing reactions between two CoA thioesters, with malonyl CoA as one subtrate. condensing reactions are similar to the proposed condensing 20 reaction for the cytoplasmic membrane bound elongase enzymes, in that in both cases an enzyme condenses two CoA thioester molecules to form two products: a ß-ketoacyl-CoA thioester and a carbon dioxide. The region of homology between the jojoba gene and chalcone synthase includes the 25 chalcone synthase active site (Lanz et al. "Site-directed mutagenesis of reservatrol and chalcone synthase, two key enzymes in different plant specific pathways" (1991) J. Biol. Chem., 266:9971-6). This active site is postulated to be involved in forming an enzyme-fatty acid 30 intermediate.

Homology was also detected between the jojoba protein and KASIII. KASIII is a soluble enzyme which catalyzes the condensation of a CoA thioester to an ACP thioester, resulting in a ß-ketoacyl-ACP thioester. A carbon dioxide molecule is released in this reaction.

While not concusive, these noted homologies suggest that the jojoba enzyme may have ß-ketoacyl-CoA synthase activity.

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Example 12 - Analysis of Plants By a fi-Keto-acyl-CoA Synthase Assay

A. The activity of ß-Keto-acyl-CoA synthase may be directly assayed in plants according to the following procedure.

Developing seeds are harvested after pollination and frozen at -70°C. For Brassica napus, the seeds are harvested 29 days after pollination. An appropriate number of seeds are thawed and homogenized in 1 ml 50 mM Hepes-NaOH, pH 7.5, 2 mM EDTA, 250 mM NaCl, 5 mM b-mercaptoethanol (twenty seeds per assay for Brassica napus). The homogenate is centrifuged at 15,000 X g for 10 min, and the oil layer is discarded. The supernatant fraction is collected and centifuged again at 200,000 X g for 1 hour.

The pellet is then resuspended in 1 ml of homogenization buffer and centrifuged a second time at 200,000 X g for 1 hour. The pellet is resuspended in 50 µl of 100 mM Hepes-NaOH, pH 7.5, 4 mM EDTA, 10% (w/v) glycerol, 2 mM b-mercaptoethanol. 10 µl of the sample is added to 10 µl of a reaction mixture cocktail and incubated at 30°C for 15 min. The final concentrations of components in the reaction mixture are: 100 mM Hepes-NaOH, pH 7.5, 1 mM b-mercaptoethanol, 100 mM oleyl CoA, 44 µM [2-14C] malonyl CoA, 4 mM EDTA and 5% (w/v) glycerol.

The reaction is stopped and the ß-ketoacyl product reduced to a diol by adding 400 μl of reducing agent solution comprised of 0.1 M K2HPO4, 0.4 M KCl, 30 % (v/v) tetrahydrofuran, and 5 mg/ml NaBH4 (added to the solution just prior to use). The mixture is incubated at 37°C for 30 min. Neutral lipids are extracted from the sample by addition of 400 μl of toluene. Radioactivity present in 100 μl of the organic phase is determined by liquid scintillation counting. The remaining toluene extract is collected and spotted onto a silica G TLC plate. The TLC plate is developed in diethyl ether:concentrated NH4OH (100:1, v/v). The migration of the diol product of the

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reduction reaction is located by use of a cold diol standard.

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B. Using this procedure plants can be assayed to determine the level of, or lack of, detectable ß-ketoacyl synthase activity. For example, HEAR plants have high levels of ß-ketoacyl synthase activity, while canola plants do not show appreciable enzyme activity. By this assay, plant species or varieties can be screened for ß-ketoacyl synthase activity to determine candidates for transformation with the sequences of this invention to achieve altered VLCFA production, or to determine canditates for screening with probes for related enzymes.

The ß-ketoacyl-CoA synthase enzyme assays demonstrate that developing embryos from high erucic acid rapeseed contain ß-ketoacyl-CoA synthase activity, while LEAR embryos do not. Embryos from transgenic plants transformed with the jojoba cDNA exhibit restored ß-ketoacyl-CoA synthase activity.

The jojoba cDNA encoding sequence thus appears to complement the mutation that differenitiates high and low 20 erucic acid rapeseed cultivars. The phenotype of the transgenic plants transformed with the jojoba gene show that a single enzyme can catalyze the formation of 20, 22 and 24 carbon fatty acids. The seed oil from the primary LEAR transformants also contains higher levels of 22:1 than 25 20:1 fatty acids. This was also true for the majority of the individual T2 seed analyzed from the 7626-212/86-2 plant. Five T2 seeds that exhibited the highest VLCFA content also contain higher levels of 22:1 than 20:1. This suggests that the ß-ketoacyl-CoA synthase is a rate 30 limiting step in the formation of VLCFA's, and that as the enzyme activity increases in developing embryos, the fatty acid profile can be switched to the longer chain lengths. The increase in the amount of 24:1 fatty acid in the oil of transgenic HEAR plants and the increase in the amount of 35 22:1 in transgenic arabidopsis plants without a concomitant increase in the quantity of VLCFAs may be a result of a difference in substrate specificities of the jojoba, Arabidopsis, and Brassica enzymes rather than an increase

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in enzyme activity which is already abundant in HEAR and Arabidopsis.

Example 13 - Other &-Keto-acyl-CoA Synthases

The active ß-ketoacyl CoA synthase chromatographs on superose with a size consistant with the enzyme being composed of two 138 kDa subunits. This suggests that the enzyme is active as a multimer, although the enzyme may be a homodimer, a heterodimer, or a higher order multimer.

The mass of one of the subunits is estimated to be $57\ kDa$ 10 by SDS gel electrophoresis and 59 kDA by calculation of the theoretical mass from translation of the cDNA sequence. The analogous soluble enzymes in plant and bacterial FAS, $\mbox{\ensuremath{\mbox{G-ketoacyl-ACP}}}$ synthases, are active as dimers with $\sim 50~\mbox{\ensuremath{\mbox{kDa}}}$ subunits. Chalcone and Stilbene synthases are also active 15 as dimers.

The jojoba ß-ketoacyl-CoA synthase subunit is a discrete 59 kDa protein. Thus, seed lipid FAE in jojobas is comprised of individual polypeptides with discrete enzyme activities similar to a type II FAS, rather than being catalyzed by the large multifunctional proteins found in type I FAS. Since the jojoba enzyme complements a Brassica mutation in FAE, it is possible that Brassica FAE is a type I system.

25 The dBEST data bank was searched with the jojoba ßketoacyl-CoA synthase DNA sequence at the NCBI using BLAST software (Altschul et al., 1990). Two Arabidopsis clones (Genbank accession Z26005, Locus 39823; and genbank accession TO4090, Locus315250) homologous to the jojoba CE cDNA were detected. The 39823 clone exhibited a high 30 degree of homology with the jojoba &-ketoacyl-CoA synthase clone. PCR primers were designed to PCR amplify and clone this sequence from Arabidopsis genomic DNA. No mRNA was detected in either developing Arabidopsis or developing 35 Brassica seeds that cross hybridized with this clone. The probe was also hybridized to RFLP blots designed to determine if homologous sequences segregate with the difference between HEAR and LEAR lines. At low hybridization stringency too many cross hybridizing bands

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are present to detect polymorphism between the HEAR and LEAR lines. At higher hybridization stringency, the bands did not cosegregate with the HEAR phenotype.

In order to isolate clones that encode related

enzymes, the protein sequences of the jojoba &-ketoacyl-CoA synthase and the Arabidopsis locus 398293 were compared to find conserved domains. Several peptide sequences were identical in the jojoba &-ketoacyl-CoA synthase and the translation of the Arabidopsis homologue 398293. Two

peptides: 1) NITTLG (amino acids 389 to 394 of the jojoba &-ketoacyl-CoA synthase) and 2) SNCKFG (amino acids 525 to 532 of the jojoba &-ketoacyl-CoA synthase) were also present in the translation of 398293. Degenerate oligonucleote primers AAYATHACNACNYTNGG and

15 SWRTTRCAYTTRAANCC encode the sense and antisense strands of the respective peptides.

The above primers PCR amplify an approximately 430 bp DNA fragment from both the jojoba ß-ketoacyl-CoA synthase cDNA and the Arabidopsis 398293 sequence. These primers can be used to PCR amplify DNA sequences that encode 20 related proteins from other tissues and other species that share nearly idendical amino acids at these conserved peptides. Using the degenerate oligonucleotides Arabidopsis green silique, HEAR, and LEAR RNA were subjected to RTPCR. Prominant bands of the expected size 25 were amplified from all 3 RNAs. One clone was obtained from the reston PCR reaction, and 2 clones from the 212/86 reaction, which appear to form two classes of cDNA clones, designated CE15 and CE20. The 212/86 CE15 clone encoded the entire CE protein (Figure 5). The protein sequences 30 translated from these clones are >98% identical to one another. The clones are approximately 50% homologous to the jojoba ß-ketoacyl-CoA synthase. The C-terminal portions of the proteins are more conserved, with the cDNAs sharing about 70% identity. Northern analysis of RNA isolated from Brassica leaf tissue and developing seed tissue showed that CE20 is highly expressed in developing seeds, and is expressed at very low levels in leaves. CE15 is expressed at high levels in leaves, and at a much lower

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level in developing seeds. The CE20 class is thus most likely to be the active condensing enzyme involved in fatty acid elongation in developing *Brassica* seeds.

The original 212/86 CE20 clone was short, and did not contain the initiator methionine. The HEAR Brassica campestris library screened with the CE15 and CE20 probes was of poor quality, and yielded only short clones. Thus, 5' RACE was used to clone the 5' end of the CE20 cDNA from 212/86 and from Reston. The sequence of the 5' race clones showed that coding region of CE 20 in both reston (HEAR) and 212/86 (LEAR) extended 3 amino acids past the 5' end of the 212/86 CE20 clone.

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CE20 primers were then chosen to get full-length CE20 sequences. Consequently,

CAUCAUCAUCAUGTCGACAAAATGACGTCCATTAACGTAAAG and
CUACUACUACUAGTCGACGGATCCTATTTGGAAGCTTTGACATTGTTTAG were
utilized. These are homologous to the 5' and 3' ends of
the protein coding region of CE20, respectively. These
primers were used to PCR the entire coding region of the

CE20 cDNA (by RTPCR) from 212/86 (Figure 6) and Reston
(Figure 7). Sequences were additionally designed for the
ends of the primers which facilitated cloning of the PCR
products in the CloneAmp vector (BRL), and restriction
enzyme sites were introduced to allow introduction of the
CE20 clones into the napin expression cassette for both
sense and antisense expression of CE20 in transgenic
Brassica plants.

The proteins deduced from *Brassica* clones CE15 and CE20 can be aligned with the protein sequence of the jojoba ß-ketoacyl-CoA synthase and *Arabidopsis* loci 398293 and 315250, with several regions of conserved protein sequence detectable. Different pairs of sense and antisense primers can thus be used to PCR amplify and isolate DNA encoding related ß-ketoacyl-CoA synthases from many different tissues, of both plant and animal species.

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Table 8

The CE15, and CE20 Brassica cDNA sequences shown in Figures 8, 9 and 10 and the condensing enzyme encoding sequence from jojoba (Figure 3) were used in determining the following primers from conserved amino acids.

SENSE PRIMER TO PEPTIDE KL(L/G)YHY

10 5381-CAUCAUCAUCAUGAATTCAAGCTTAARYTNBKNTAYCAYTA

SENSE PRIMER TO PEPTIDE NLGGMGC

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5384-CAUCAUCAUCAUGAATTCAAGCTTAAYYTNGGNGGNATGGG

20

ANTISENSESENSE PRIMER TO PEPTIDE NLGGMGC

5382-CUACUACUAGGATCCGTCGACCCATNCCNCCNARRTT

25

ANTISENSESENSE PRIMER TO PEPTIDE GFKCNS

5385-CUACUACUAGGATCCGTCGACSWRTTRCAYTTRAANCC

30

ANTISENSESENSE PRIMER TO PEPTIDE GFKCNS

35 4872-CUACUACUASWRTTRCAYTTRAANCC

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These primers from Table 8 were variously used to PCR (RTPCR) amplify fragments from RNA isolated from developing seeds of Lunaria annua, Tropaoelu majus (Nasturtium), and green siliques of Arabidopsis thaliana. The primers most successfully utilized were 5381-CAUCAUCAUCAUGAATTCAAGCTTAARYTNBKNTAYCAYTA (a sense primer to peptide KL(L/G)YHY) and

CUACUACUAGGATCCGTCGACCCATNCCNCCNARRTT (an antisense primer to peptide NLGGMGC). These primers were used to produce three clones encoding a portion of the elongase condensing enzyme from *Arabidopsis*, designated ARAB CE15, ARAB CE17 and ARAB CE19 (Figures 8, 9 and 10, respectively)

From Lunaria a single clone was identified, LUN CE8

(Figure 11). Since Lunaria produces high levels of 24:1 fatty acid in its seed oil (up to 30%), a cDNA library from RNA isolated from developing seeds of Lunaria was constructed, and LUN CE8 was used to screen this Lunaria cDNA library.

Three classes of cDNA clones were isolated, Lunaria 1, Lunaria 5, and Lunaria 27 (Figures 12, 13 and 14, respectively). Of total clones, 81% (26/32) of the clones isolated were of a class similar to Lunaria 5. Of the remainder, 16% (5/32) of the clones were similar to the PCR probe, LUN CE8, designated Lunaria 1. One clone, Lunaria 27, was unique.

As seen in Table 9, Lunaria 5 shares approximately 85% homology with the Brassica CE20 clones. The high degree of homology with the Brassica seed expressed cDNA, and the high abundance of the Lunaria 5 cDNA in developing seed tissue suggest that Lunaria 5 is the cDNA that is active in seed oil fatty acid elongation.

Table 9

Sequence pair distances based on the BIG ALIGNM program, using a Clustal method with PAM250 residue weight table.

5

Percent Divergence

Per	cent	Simi	larity

				1 6106	311L OIII	manty			
Į		1	2	3	4	5	6	7	
	1		55.6	55.4	53.0	51.2	59.0	67.9	1
	_2	44.7		99.1	85.1	41.0	61.7	52.3	2
	3	43.5	0.7		85.2	40.6	61.7	52.8	3
	4	44.7	16.1	16.2		40.5	63.4	53.0	4
ı	_5	44.8	53.1	53.1	52.5		49.1	49.1	5
i	- 6	40.6	37.9	38.9	36.4	43.7		58.8	6
	7	33.0	45.6	46.0	45.0	46.3	39.2		7
		1	2	3	4	5	6	7	·

JOJOBA
212/86 CE20
RESTON CE20
LUNARIA 5 (PRELIMINARY)
212/86 CE15
LUNARIA 1 (PRELI
LUNARIA 27 (PREL

SUBSTITUTE SHEET (RULE 26)

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Finally, a partial *Nasturtium* PCR clone was obtained using the same primers as were used to isolate LUN CE8. The sequence to the nasturtuim clone (NAST CE26) is provided in Figure 15.

The use of ß-ketoacyl-CoA synthases obtained in this manner from other tissues or other species that have different substrate specificities can be used to create modified seed oils with different chain length fatty acids. This could include enzymes isolated from plant taxa such as lunaria, which synthesizes significant quantities of 24:1 fatty acid in its seed tissue. This could also include enzymes involved in cuticular wax synthesis of any plant species which may be capable of synthesizing fatty acids of chain lengths greater than 24 carbons. For instance, 15 Lunaria seeds contain up to 30% 24:1 in their seed oil. Condensing enzyme assay on crude extract from developing Lunaria seeds shows that the enzyme is active at elongating 18:1 to 20:1, 20:1 to 22:1 and 22:1 to 24:1. These data 20 suggest that the Lunaria enzyme will be useful for producing 24:1 in transgenic plants. As it is, expression of the jojoba enzyme in transgenic Brassica has resulted in plants having up to 7.8% of the seed oil composed of 24:1. The source jojoba seeds only produce 4.1 % of the oil in the seed as 24:1. The above respresents the first 25 description of an approach for increasing the 24:1 content of transgenic oil.

The above Examples also demonstrate that the primers of Table 7 can be used to successfully isolate condensing enzyme clones from diverse plant species. These oligonucleotides may be especially useful for isolating the corresponding fatty acid synthase animal genes, which have not been previously cloned. Since the ß-ketoacyl-CoA synthase expression is repressed in several demyelinating nervous system disorders of humans, for instance adrenoleukodystrophy, adrenomyeloneuropathy, and multiple svlrtodid(reviewed in Sargent and Coupland, 1994), the human genes may be useful in human gene therapy.

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Similarly, vegetable oils high in 22:1 or 24:1 may be useful dietary therapeutic agents for these diseases.

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All publications and patent applications cited in this specification are herein incorporated by reference as if each individual publication or patent application were specifically and individually indicated to be incorporated by reference.

Although the foregoing invention has been described in some detail by way of illustration and example for purposes of clarity and understanding, it will be readily apparent to those of ordinary skill in the art in light of the teaching of this invention that certain changes and modifications may be made thereto without departing from the spirit or scope of the appended claims.

CLAIMS

What is claimed is:

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1. A method for the production of a 24:1 very long chain fatty acid molecule in a plant seed cell, said plant otherwise incapable of producing seed having more than 5% by weight of said very long chain fatty acid molecule, said method comprising the steps of:

growing a plant under conditions wherein said plant produces long chain fatty acyl-CoA molecules in the plant seed, in the presence of an expression product of a very long chain fatty acid molecule-altering DNA sequence operably linked to regulatory elements for directing the expression of said DNA sequence such as to effect the contact between such long chain fatty acyl-CoA molecules and said expression product, and producing said very long chain fatty acid molecule in said plant seed at a level above 5% by weight.

2. The method of Claim 1 wherein said very long chain fatty acid molecule is produced in said plant seed to a level greater than 7% by weight.

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3. The method of Claim 1 wherein said regulatory elements direct preferrential expression of said DNA sequence in plant seed embryo cells.

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- 4. The method of Claim 1 wherein said very long chain fatty acid molecule-altering DNA sequence is a condensing enzyme encoding sequence from *Brassica*.
- 5. The method of Claim 4 wherein said *Brassica* sencoding sequence is to the CE15 class of condensing enzymes.

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- 6. The method of Claim 4 wherein said Brassica encoding sequence is to the CE20 class of condensing enzymes.
- 5 7. The method of Claim 1 wherein said very long chain fatty acid molecule-altering DNA sequence is a condensing enzyme encoding sequence from Arabadopsis.
- 8. The method of Claim 1 wherein said very long chain 10 fatty acid molecule-altering DNA sequence is a condensing enzyme encoding sequence from Nasturtium.
- 9. The method of Claim 1 wherein said very long chain fatty acid molecule-altering DNA sequence is a condensing enzyme encoding sequence from *Lunaria*.
 - 10. The method of Claim 9 wherein said Lunaria encoding sequence is Lunaria 5.
- 20 11. The method of Claim 1 wherein said regulatory elements direct preferrential expression of said DNA sequence in plant seed embryo cells.
- 12. A plant seed containing a very long chain fatty 25 acid molecule produced in accordance with Claim 1.
 - 13. A plant seed produced in accordance with Claim 1.
- 14. A method for decreasing the proportion of VLCFA 30 in a plant from a given proportion of VLCFA comprising the steps of:

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growing a plant under conditions wherein said plant produces VLCFA and ß-ketoacyl-CoA synthase, in the presence of a ß-ketoacyl-CoA-decreasing DNA sequence operably linked to regulatory elements for directing the expression of said DNA sequence in said cell, wherein said DNA sequence encodes a ß-ketoacyl-CoA DNA sequence of said plant and the expression of said DNA sequence results in a decrease in the production of ß-ketoacyl-CoA synthase by said plant

cell and a decrease in the proportion of VLCFA produced by said plant cell.

- 15. The method of Claim 14 wherein said regulatory elements direct the antisense transcription of said DNA sequence.
- 16. The method of Claim 14 wherein said regulatory elements direct preferrential expression of said DNA

 10 sequence in plant seed embryo cells and wherein said VLCFA and said ß-keto acyl-CoA is produced in plant seed.
 - 17. A plant seed cell produced in accordance with Claim 9.

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- 18. A construct comprising a DNA sequence which encodes a condensing enzyme and a heterologous DNA sequence not naturally associated with said encoding sequence wherein said condensing enzyme encoding sequence is 20 obtained by screening a DNA library prepared from an organism which is capable of producing very long chain fatty acid molecules with degenerate oligonucleotide primers selected from the group consisting of CAUCAUCAUCAUGAATTCAAGCTTAARYTNBKNTAYCAYTA,

 25 CAUCAUCAUGAATTCAAGCTTAAYYTNGGNGGNATGGG, CUACUACUACUAGGATCCGTCGACCCATNCCNCCNARRTT, CUACUACUACUAGGATCCGTCGACSWRTTRCAYTTRAANCC and
- 19. An isolated nucleic acid sequence encoding a condensing enzyme which can be isolated according to a method comprising the step of PCR amplification utilizing primers CAUCAUCAUCAUGAATTCAAGCTTAARYTNBKNTAYCAYTA and CUACUACUACUAGGATCCGTCGACCCATNCCNCCNARRTT.

CUACUACUASWRTTRCAYTTRAANCC.

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20. A construct comprising a nucleic sequence according to Claim 19 and a heterologous DNA sequence not naturally associated with said encoding sequence.

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21. A construct according to Claim 20 wherein said heterologous DNA sequence comprises regulatory elements which direct preferrential expression of said DNA sequence in plant seed embryo cells.

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- 22. A construct according to Claim 20 wherein said condensing enzyme encoding sequence is from Brassica.
- A construct according to Claim 22 wherein said Brassica encoding sequence is to the CE15 class of 10 condensing enzymes.
- 24. A construct according to Claim 22 wherein said Brassica encoding sequence is to the CE20 class of 15 condensing enzymes.
 - 25. A construct according to Claim 20 wherein said condensing enzyme encoding sequence is from Arabadopsis.
- 20 26. A construct according to Claim 20 wherein said condensing enzyme encoding sequence is from Nasturtium.
 - 27. A construct according to Claim 20 wherein said condensing enzyme encoding sequence is from Lunaria.

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28. A construct according to Claim 27 wherein said Lunaria encoding sequence is Lunaria 5.

					•
09	112	160	208	256	304
AAATCCTCCA CTCATACACT CCACTTCTCT CTCTCTCT	GTAGCAAACT TAAAAGAAA ATG GAA ATG GGA AGC ATT TTA GAG TTT CTT Met Glu Glu Met Gly Ser Ile Leu Glu Phe Leu 1	GAT AAC AAA GCC ATT TTG GTC ACT GGT GCT ACT GGC TCC TTA GCA AAA Asp Asn Lys Ala Ile Leu Val Thr Gly Ala Thr Gly Ser Leu Ala Lys 15	ATT TTT GTG GAG AAG GTA CTG AGG AGT CAA CCG AAT GTG AAG AAA CTC Ile Phe Val Glu Lys Val Leu Arg Ser Gln Pro Asn Val Lys Lys Leu 30	TAT CTT/CTT TTG AGA GCA ACC GAT GAC GAG ACA GCT GCT CTA CGC TTG Tyr Leu Leu Leu Arg Ala Thr Asp Asp Glu Thr Ala Ala Leu Arg Leu 50	CAA AAT GAG GTT TTT GGA AAA GAG TTG TTC AAA GTT CTG AAA CAA AAT Gln Asn Glu Val Phe Gly Lys Glu Leu Phe Lys Val Leu Lys Gln Asn 60 75

FIG. 1A

				•	
352	400	448	496	544	592
ITA GGT GCA AAT TTC TAT TCC TTT GTA TCA GAA AAA GTG ACT GTA GTA Leu Gly Ala Asn Phe Tyr Ser Phe Val Ser Glu Lys Val Thr Val Val 80	CCC GGT GAT ATT ACT GGT GAA GAC TTG TGT CTC AAA GAC GTC AAT TTG Pro Gly Asp Ile Thr Gly Glu Asp Leu Cys Leu Lys Asp Val Asn Leu 95	AAG GAA GAA ATG TGG AGG GAA ATC GAT GTT GTT GTC AAT CTA GCT GCT Lys Glu Glu Met Trp Arg Glu Ile Asp Val Val Val Asn Leu Ala Ala 110	ACA ATC AAC TTC ATT GAA AGG TAC GAC GTG TCT CTG CTT ATC AAC ACA Thr Ile Asn Phe Ile Glu Arg Tyr Asp Val Ser Leu Leu Ile Asn Thr 125	TAT GGA GCC AAG TAT GTT TTG GAC TTC GCG AAG AAG TGC AAC AAA TTA Tyr Gly Ala Lys Tyr Val Leu Asp Phe Ala Lys Lys Cys Asn Lys Leu 140	AAG ATA TTT GTT CAT GTA TCT ACT GCT TAT GTA TCT GGA GAG AAA AAT Lys Ile Phe Val His Val Ser Thr Ala Tyr Val Ser Gly Glu Lys Asn 160
		,* * H	7	HHH	4 J

640	889	736	784	832	880
GGC GAG TCA CTT AAT GGA	AAA CTT GTG GAG GCA AAA	GAA AAG TCC ATT AAA TCG	3A CAC TGG GGA TGG CCA	NG ATG CTT TTG ATG CAA	T CCC ACC ATC ATC ACC
Gly Glu Ser Leu Asn Gly	Lys Leu Val Glu Ala Lys	Glu Lys Ser Ile Lys Ser	cg His Trp Gly Trp Pro	.u Met Leu Leu Met Gln	G Pro Thr Ile Ile Thr
185	200	215	30	250	265
GAG AAG CCT TAT TAT ATG G	GAC ATT AAT GTA GAG AAG AA	CAA GCA GCG GGG GCA ACG GA	ATG GGC ATC GAG AGG GCA AGA	TTC ACC AAG GCA TTA GGG GAG	ATT CCG CTT ACT ATT ATT CGT
Glu Lys Pro Tyr Tyr Met G	Asp Ile Asn Val Glu Lys L ₃	Gln Ala Ala Gly Ala Thr Gl	Met Gly Ile Glu Arg Ala Arg	Phe Thr Lys Ala Leu Gly Glu	Ile Pro Leu Thr Ile Ile Arg
180	195	210	225	240	260
GGG TTA ATA CTG (Gly Leu Ile Leu (175	AGA TTA GGT CTG G Arg Leu Gly Leu A	ATC AAT GAA CTT C Ile Asn Glu Leu G 205	ACA ATG AAG GAC A Thr Met Lys Asp M 220	AAT GTG TAT GTA T' Asn Val Tyr Val Pl 24	TAC AAA GGG GAC AI Tyr Lys Gly Asp II 255

928	976	1024	1072	1120	1168
ACC	ATG Met	GTC Val 315	TAC	ATG Met	AAT Asn
AGG Arg	TGT Cys	ATG Met	AGA Arg 330	CCA Pro I	AAG 1 Lys 1
GTC Val	AGG Arg	GAT	CAA	AAT (Asn) 345	ACC 7 Thr 1
GGT G1y 280	TTG Leu	GCA	AAC Asn (GCG A	TTC 7 Phe 1 360
GAA Glu	AGA Arg 295	CCG	GCA Ala	GCG Ala	TAC TYT I
GTT Val	GGG G1y	ATA Ile 310	CAC His	Ser	CGT /
TGG Trp	AAA Lys	CTG	GCG Ala 325	TCT	CAC (
GGT Gly	GGT G1y	GAC Asp	GTG Val	GGA G1y 340	GCA (
CCT Pro 275	TAT Tyr	ATT Ile	ATG Met	GTG Val	ATG (Met 7
rrr Phe	TAT Tyr 290	ATA Ile	GCC	CAT (GAG Z
CCC	GTA Val	ACA Thr 305	GTA	TAC TYT)	CCA (Pro (
GAG Glu	CCT	AGC Ser	ATA Ile 320	ACA Thr	TTA
AAA Lys	GTA Val	CCC	ACG Thr	GTG Val 335	GCA Ala
rrr Phe 270	AAT Asn	GGA Gly	GCA Ala	CCG	AGT (Ser 350
ACT Thr	GAT Asp 285	TGC	AAT	GAG Glu	CTG
AGC	ATC Ile	CTT Leu 300	GTG Val	GTA Val	AAA Lys

1216	1264	1312	1360	1408	1456
TGG ATC AAC CCG GAT CGC AAC CCA GTA CAT GTG GGT CGG GCT ATG Trp Ile Asn Pro Asp Arg Asn Pro Val His Val Gly Arg Ala Met 365	TTC TCC TCC TTC TCC ACC TTC CAC CTT TAT CTC ACC CTT AAT TTC Phe Ser Ser Phe Ser Thr Phe His Leu Tyr Leu Thr Leu Asn Phe 385	CTT CCT TTG AAG GTA CTG GAG ATA GCA AAT ACA ATA TTC TGC CAA Leu Pro Leu Lys Val Leu Glu Ile Ala Asn Thr Ile Phe Cys Gln 400	TTC AAG GGT AAG TAC ATG GAT CTT AAA AGG AAG ACG AGG TTG TTG 1 Phe Lys Gly Lys Tyr Met Asp Leu Lys Arg Lys Thr Arg Leu Leu 415	CGT TTA GTA GAC ATT TAT AAA CCC TAC CTC TTC TTC CAA GGC ATC 1. Arg Leu Val Asp Ile Tyr Lys Pro Tyr Leu Phe Phe Gln Gly Ile 430	GAT GAC ATG AAC ACT GAG AAG TTG CGG ATT GCT GCA AAA GAA AGC 14 Asp Asp Met Asn Thr Glu Lys Leu Arg Ile Ala Ala Lys Glu Ser 445
CCA	GTC Val 380	CTC	TGG	TTG (Len)	TTT (Phe 1

1504	1552	1608	1668	1728	1786
ATA GTT GAA GCT GAT ATG TTT TAC TTT GAT CCC AGG GCA ATT AAC TGG Ile Val Glu Ala Asp Met Phe Tyr Phe Asp Pro Arg Ala Ile Asn Trp 460	GAA GAT TAC TTC TAG AAA ACT CAT TTC CCA GGN GTC GTA GAG CAC GTT Glu Asp Tyr Phe Leu Lys Thr His Phe Pro Gly Val Val Glu His Val 490	CTT AAC TAAAAGTTAC GGTACGAAAA TGAGAAGATT GGAATGCATG CACCGAAAGN Leu Asn	NCAACATAAA AGACGTGGTT AAAGTCATGG TCAAAAAAGA AATAAAATGC AGTTAGGTTT 1	GIGITGCAGT TITGAITCCT IGIAITGITA CITGTACTIT IGALCTITIT CTITITIAAI 1728	GAAATTTCTC TCTTTGTTTT GTGAAAAAA AAAAAAAAA GAGCTCCTGC AGAAGCTT 1

					•
56	104	152	200	248	296
GGAACTCCAT CCCTTCCTCC CTCACTCCTC TCTCTACA ATG AAG GCC AAA ACA ATC Met Lys Ala Lys Thr Ile 1	ACA AAC CCG GAG ATC CAA GTC TCC ACG ACC ATG ACC ACC ACG ACC ACG Thr Abr Thr Thr Thr Thr Thr Thr Thr Thr Thr 10	ACT ATG ACC GCC ACT CTC CCC AAC TTC AAG TCC TCC ATC AAC TTA CAC Thr Met Thr Ala Thr Leu Pro Asn Phe Lys Ser Ser Ile Asn Leu His 25	CAC GTC AAG CTC GGC TAC CAC TTA ATC TCC AAT GCC CTC TTC CTC His Val Lys Leu Gly Tyr His Tyr Leu Ile Ser Asn Ala Leu Phe Leu 40	GTA TTC ATC CCC CTT TTG GGC CTC GCT TCG GCC CAT CTC TCC TCC TTC Val Phe Ile Pro Leu Leu Gly Leu Ala Ser Ala His Leu Ser Ser Phe 55 65	TCG GCC CAT GAC TTG TCC CTG CTC TTC GAC CTC CTT CGC CGC AAC CTC Ser Ala His Asp Leu Ser Leu Leu Phe Asp Leu Leu Arg Arg Asn Leu 75

FIG. 2A

T GTT GTC GTT TGT TCT TTC CTC TTC GTT TTA TTR Val Val Val Val Cys Ser Phe Leu Phe Val Leu Leu 95 TTG ACC CGG CCC AGG AAT GTC TAC TTG GTG GAC Leu Thr Arg Pro Arg Asn Val Tyr Leu Val Asp 115 115 CCT CAA CCG AAC CTG ATG ACA TCC CAC GAG ATG 125 Pro Gln Pro Asn Leu Met Thr Ser His Glu Met 125 TCC CGG GCC GGG TCG TTT TCT AAG GAG AAT ATT Ser Arg Ala Gly Ser Phe Ser Lys Glu Asn Ile 145 ATC TTG GAG AGG GCC GGT ATG GGT CGG GAA ACC Ile Leu Glu Arg Ala Gly Met Gly Arg Glu Thr 155 GTC ACT AAG GTG CCC GAG CCG AGC ATA GCA Val Thr Lys Val Pro Ala Glu Pro Ser Ile Ala 170	344	392	440	488	536	584
CTC CC Leu Procession Procession CCG ACC ACC ACC ACC ACC ACC ACC ACC ACC	CCT GTT GTC GTT TGT TCT TTC CTC TTC GTT TTA TTA GCA ACC Pro Val Val Val Cys Ser Phe Leu Phe Val Leu Leu Ala Thr 90	TTC TTG ACC CGG CCC AGG AAT GTC TAC TTG GTG GAC TTT GGA Phe Leu Thr Arg Pro Arg Asn Val Tyr Leu Val Asp Phe Gly 105	AAG CCT CAA CCG AAC CTG ATG ACA TCC CAC GAG ATG TTC ATG. Lys Pro Gln Pro Asn Leu Met Thr Ser His Glu Met Phe Met 120	ACC TCC CGG GCC GGG TCG TTT TCT AAG GAG AAT ATT GAG TTT Thr Ser Arg Ala Gly Ser Phe Ser Lys Glu Asn Ile Glu Phe 140	AAG ATC TTG GAG AGG GCC GGT ATG GGT CGG GAA ACC TAT GTC Lys lle Leu Glu Arg Ala Gly Met Gly Arg Glu Thr Tyr Val 160	TCC GTC ACT AAG GTG CCC GCC GAG CCG AGC ATA GCA GCC Ser Val Thr Lys Val Pro Ala Glu Pro Ser Ile Ala Ala 170

				•	
632	089	728	776	824	872
GTG TTG GAG Val Leu Glu	GTG ANC TGC Val Xxx Cys	GTT AAC CAT Val Asn His 230	GGC ATG GGT Gly Met Gly 245	CTC CTA CAG Leu Leu Gln 260	GAA AAC ATG Glu Asn Met
GAG Glu 195	GTG Val	ATA Ile	GGT Gly	GAC Asp	ACG
c GAC e Asp	A CTG P Leu 210	C ATG	CTT Leu	AAG Lys	AGC Ser
G ATC a Ile	A ATA Y Ile	A TCC r Ser 225	r AAT r Asn 0	r GCC ı Ala	A GTG L Val
GGG GCG Gly Ala	ATA GGA Ile Gly	CTG TCA Leu Ser	AGC TAT Ser Tyr 240	AT CTT Sp Leu is	'A GTA 'u Val
TAC G TYr G 190	CAG A	TCG CT Ser Le	CTT AC Leu Se	ATT GAT Ile Asp 255	GTG TTA Val Leu
ATG	AAG Lys 205	CCG	ATA Ile	TCC Ser	TAT (Tyr 1
GTG 1 Val	G CCG	ACG Thr 220	AAT	ATT Ile	ACA
G GAG u Glu	G AAG 1 Lys	c ccA n Pro	G GGT g G1Y 235	3 CTC Y Leu J	A AAC s Asn
GCG GAG Ala Glu 185	GGG GTG Gly Val	TYY AAC Phe Asn	CTN AGG Leu Arg	T GGG .a Gly 250	T AAA G Lys
GAG GG Glu A	ACG G(Thr G] 200	TTG TY Leu Pł	AAG CT Lys Le	AGT GCT Ser Ala	TAC CGT Tyr Arg
GCC (Ala (AAG A Lys 1	AGC 1 Ser I 215	TAC A Tyr I	TGC A Cys S	GTT T Val T

FIG. 2C

920	896	1016	1064	1112	1160	1208
CTT AAT TGG TAC TGG GGC AAT GAC CGC TCC ATG CTT ATC ACC Leu Asn Trp Tyr Trp Gly Asn Asp Arg Ser Met Leu Ile Thr Asn 280	CTA TTT CGC ATG GGT GGC GCT GCC ATC ATC CTC TCA AAC CGC TGG Leu Phe Arg Met Gly Gly Ala Ala Ile Ile Leu Ser Asn Arg Trp 300	GAT CGT CGC CGA TCC AAG TAC CAA CTC CTT CAT ACA GTA CGC ACC ASP Asp Arg Arg Ser Lys Tyr Gln Leu Leu His Thr Val Arg Thr 320	AAG GGC GCT GAC GAC AAG TCC TAT AGA TGC GTC TTA CAA CAA GAA Lys Gly Ala Asp Asp Lys Ser Tyr Arg Cys Val Leu Gln Gln Glu 330	GAA AAT AAC AAG GTA GGT GTT GCC TTA TCC AAG GAT CTG ATG GCA 11 Glu Asn Asn Lys Val Gly Val Ala Leu Ser Lys Asp Leu Met Ala 345	GCC GGT GAA GCC CTA AAG GCC AAC ATC ACG ACC CTT GGT CCC CTC 11 Ala Gly Glu Ala Leu Lys Ala Asn Ile Thr Thr Leu Gly Pro Leu 360	CTC CCC ATG TCA GAA CAA CTC CTC TTC TTT GCC ACC TTA GTG GCA 12 Leu Pro Met Ser Glu Gln Leu Leu Phe Phe Ala Thr Leu Val Ala 380
ACC	TGC Cys 295	CGT	CAC His	GAT Asp	GTT Val	GTG Val 375

1256	1304	1352	1400	1448	1496
CGT AAG GTC TTC AAG ATG ACG AAC GTG AAG CCA TAC ATC CCA GAT TTC 1 Arg Lys Val Phe Lys Met Thr Asn Val Lys Pro Tyr Ile Pro Asp Phe 395	TTG GCA GCG AAC GAC TTC TGC ATC CAT GCA GGA GGC AAA GCA GTG Leu Ala Ala Asn Asp Phe Cys Ile His Ala Gly Gly Lys Ala Val 410	GAT GAG CTC GAG AAG AAC TTG GAG TTG ACG CCA TGG CAC CTT GAA Asp Glu Leu Glu Lys Asn Leu Glu Leu Thr Pro Trp His Leu Glu 425	TCG AGG ATG ACA CTG TAT AGG TTT GGG AAC ACA TCG AGT AGC TCA Ser Arg Met Thr Leu Tyr Arg Phe Gly Asn Thr Ser Ser Ser 440	TGG TAC GAG TTG GCA TAC GCT GAA GCA AAA GGG AGG ATC CGT AAG Trp Tyr Glu Leu Ala Tyr Ala Glu Ala Lys Gly Arg Ile Arg Lys 460	GAT CGA ACT TGG ATG ATT GGA TTT GGT TCA GGT TTC AAG TGT AAC ASP Arg Thr Trp Met Ile Gly Phe Gly Ser Gly Phe Lys Cys Asn 475
υ«	AAG Lys	TTG	CCC	TTA Leu 455	$_{\rm G1y}^{\rm GGT}$

FIG. 2E

1544	1592	1640	00	33
11	45	16	17	1733
AGT GTT GTG AGG GCT TTG AGG AGT GTC AAT CCG GCT AGA GAG AAG Ser Val Val Trp Arg Ala Leu Arg Ser Val Asn Pro Ala Arg Glu Lys 495	AAT CCT TGG ATG GAA ATT GAG AAG TTC CCT GTC CAT GTG CCT AAA Asn Pro Trp Met Asp Glu Ile Glu Lys Phe Pro Val His Val Pro Lys 505	ATC GCA CCT ATC GCT TCG TAGAACTGCT AGGATGTGAT TAGTAATGAA Ile Ala Pro Ile Ala Ser 520	AAATGTGTAT TATGTTAGTG ATGTAGAAAA AGAAACTTTA GTTGATGGGT GAGAACATGT 1700	CTCATTGAGA ATAACGTGTG CATCGTTGTG TTG

IG. 2F

51	66	147	195	243	291
GTCGACACA ATG AAG GCC AAA ACA ATC ACA AAC CCG GAG ATC CAA GTC TCC	ACG ACC ATG ACC ACG ACC ACG ACC GCC ACT CTC CCC AAC TTC AAG	TCC TCC ATC AAC TTA CAC CAC GTC AAG CTC GGC TAC CAC TAC TTA ATC	TCC AAT GCC CTC TTC CTC GTA TTC ATC CCC CTT TTG GGC CTC GCT TCG	GCC CAC CTC TCC TTC TCG GCC CAT GAC TTG TCC CTG CTC TTC GAC	CTC CTT CGC CGC AAC CTC CTC CCC GTT GTC GTT TGT TCT TTC CTC TTC
Met Lys Ala Lys Thr Ile Thr Asn Pro Glu Ile Gln Val Ser	Thr Thr Met Thr Thr Thr Thr Thr Ala Thr Leu Pro Asn Phe Lys	Ser Ser Ile Asn Leu His His Val Lys Leu Gly Tyr His Tyr Leu Ile	Ser Asn Ala Leu Phe Leu Val Phe Ile Pro Leu Leu Gly Leu Ala Ser	Ala His Leu Ser Ser Phe Ser Ala His Asp Leu Ser Leu Leu Phe Asp	Leu Leu Arg Arg Asn Leu Leu Pro Val Val Val Cys Ser Phe Leu Phe
1 5	15	35	50 55	65	80

FIG. 3A

	٠				
339	387	435	483	531	579
TAC Tyr 110	TCC Ser	AAG Lys	GGC Gly	CCG	GCG Ala 190
GTC Val	ACA Thr 125	TCT Ser	ATG Met	GAG Glu	GGG G1y
AAT Asn	ATA Ile	TTT Phe 140	6GT Glу	CCC	TAC (Tyr (
AGG Arg	CTG	TCG Ser	GCC Ala 155	CCG	ATG Met
CCT	AAC Asn	GGG G1y	AGG Arg	GTG Val 170	GTG Val 1
CGG Arg 105	CCG Pro	GCC Ala	GAG Glu	AAG Lys	GAG Glu 185
ACC Thr	CAC His 120	CGG Arg	TTG	ACT Thr	GAG
TTG	CCT	TCC Ser 135	ATC Ile	GTC Val	GCG
TTC	AAG Lys	ACC	AAG Lys 150	Ser	GAG Glu
CAT His	TAT Tyr	CGG Arg	AGG Arg	GAA Glu 165	GCC
CTA Leu 100	TGC	GAC Asp	CAG Gln	CCC	AGG Arg 180
ACC Thr	GCC Ala 115	ATG Met	TTT Phe	GTC Val	GCC Ala
GCA Ala	TTT Phe	TTC Phe 130	GAG Glu	TAC Tyr	GCA Ala
TTA	GAC Asp	ATG Met	ATT Ile 145	ACC Thr	GCA Ala
TTA Leu	GTG Val	GAG Glu	AAT Asn	GAA Glu 160	ATA
GTT Val 95	TTG	CAC His	GAG Glu	CGG Arg	AGC Ser 175

627	675	723	771	819	867
GGA Gly	TCA Ser	TAT Tyr	CTT Leu	GTG Val 270	TCC Ser
ATA Ile 205	CTG	AGC Ser	GAT Asp	GTA	CGC Arg 285
CAG Gln	TCG Ser 220	CTT Leu	ATT Ile	TTA (GAC (
AAG Lys	CCG Pro	ATA I1e 235	TCC Ser	GTG Val	AAT (Asn i
CCG	ACG	AAT Asn	ATT Ile 250	TAT Tyr	GGC 7
AAG Lys	CCA Pro	GGT Gly	CIC	ACA Thr 265	TGG (Trp (
GTG Val 200	AAC Asn	AGG Arg	GGG Gly	AAC Asn	TAC Tyr 280
GGG G1y	TTT Phe 215	CTT	GCT Ala	CGT	TGG
ACG Thr	TTG	AAG Lys 230	AGT Ser	TAC	AAT Asn
AAG Lys	AGC	TAC	TGC Cys 245	GTT Val	CTT
GAG Glu	TGC	CAT His	GGT G1y	CAG Gln 260	ACC
TTG Leu 195	AAC Asn	AAC Asn	ATG Met	CTA	ATG Met 275
GTG Val	GTG Val 210	GTT Val	66C G1y	CTC	AAC Asn
GAG Glu	GTG Val	ATA Ile 225	GGT Gly	GAC Asp	GAA Glu
GAC ASP	CTG	ATG Met	CTT Leu 240	AAG Lys	ACA Thr
ATC Ile	ATA Ile	TCC	AAT Asn	GCC Ala 255	AGC

FIG. 3C

			1059	1107	1155
ATC ACC AAC TGC CTA TTT CGC ATG GGT GGC GCT GCC ATC ATC Ile Thr Asn Cys Leu Phe Arg Met Gly Gly Ala Ala Ile Ile 290	AAC CGC TGG CGT GAT CGT CGC CGA TCC AAG TAC CAA CTC CTT Asn Arg Trp Arg Asp Arg Arg Ser Lys Tyr Gln Leu Leu 305	GTA CGC ACC CAC AAG GGC GCT GAC GAC AAG TCC TAT AGA TGC Val Arg Thr His Lys Gly Ala Asp Asp Lys Ser Tyr Arg Cys 330	CAA CAA GAA GAT GAA AAT AAC AAG GTA GGT GTT GCC TTA TCC 1 Gln Gln Glu Asp Glu Asn Asn Lys Val Gly Val Ala Leu Ser 345	CTG ATG GCA GTT GCC GGT GAA GCC CTA AAG GCC AAC ATC ACG 1 Leu Met Ala Val Ala Gly Glu Ala Leu Lys Ala Asn Ile Thr 355	GGT CCC CTC GTG CTC CCC ATG TCA GAA CAA CTC CTC TTC TTT 1. Gly Pro Leu Val Leu Pro Met Ser Glu Gln Leu Leu Phe Phe 370
ATG CTT A Met Leu I	CTC TCA A Leu Ser A 3	CAC ACA G' His Thr Va 320	GTC TTA C2 Val Leu G] 335	AAG GAT CI Lys Asp Le	ACC CTT GG Thr Leu Gl

FIG. 3D

1203	1251	1299	1347	1395	1443
ACC TTA GTG GCA CGT AAG GTC TTC AAG ATG ACG AAC GTG AAG CCA 1.2 Thr Leu Val Ala Arg Lys Val Phe Lys Met Thr Asn Val Lys Pro 385	ATC CCA GAT TTC AAG TTG GCA GCG AAG CAC TTC TGC ATC CAT GCA 12 Ile Pro Asp Phe Lys Leu Ala Ala Lys His Phe Cys Ile His Ala 400	GGC AAA GCA GTG TTG GAT GAG CTC GAG ACG AAC TTG GAG TTG ACG 12 Gly Lys Ala Val Leu Asp Glu Leu Glu Thr Asn Leu Glu Leu Thr 420	TGG CAC CTT GAA CCC TCG AGG ATG ACA CTG TAT AGG TTT GGG AAC 13 Trp His Leu Glu Pro Ser Arg Met Thr Leu Tyr Arg Phe Gly Asn 445	AGT AGC TCA TTA TGG TAC GAG TTG GCA TAC GCT GAA GCA AAA Ser Ser Ser Leu Trp Tyr Glu Leu Ala Tyr Ala Glu Ala Lys 450	ATC CGT AAG GGT GAT CGA ACT TGG ATG ATT GGA TTT GGT TCA Ile Arg Lys Gly Asp Arg Thr Trp Met Ile Gly Phe Gly Ser 465
GCC A Ala T	TAC A Tyr I	GGA G Gly G 415	CCA TV Pro T	ACA TCG Thr Ser	GGG AGG Gly Arg

FIG. 3E

191	339	26	52	12	72	83
7	H		16	17	17	178
AAC AGT GTG TGG AGG GCT TTG AGG AGT GTC AAT 1491 Asn Ser Val Val Trp Arg Ala Leu Arg Ser Val Asn 490	AAG AAT CCT TGG ATG GAT GAA ATT GAG AAT TTC CCT 1539 Lys Asn Pro Trp Met Asp Glu Ile Glu Asn Phe Pro 500	AAA ATC GCA CCT ATC GCT TCG TAGAACTGCT AGGATGTGAT 1592 Lys Ile Ala Pro Ile Ala Ser 515	TAGTAATGAA AAATGTGTAT TATGTTAGTG ATGTAGAAAA AGAAACTTTA GTTGATGGGT 1652	CTCATTGAGA ATAACGTGTG CATCGTTGTG TTGAATTTGA ATTTGAGTAT 1712	TGGTGAAATT CTGTTAGAAT TGACGCATGA GTCATATATA TACAAATTTA AGTAAGATTT 1772	1783
TGT Cys	GAG Glu	CCT	AAT	rca:	rgiri	
AAG Lys	AGA Arg	GTG (AA A	GT C	TT CI	CT
TTC Phe 480	GCT Ala	CAT His	AATG	ACAT	3AAA	TTT
GGT GIY]	CCG (Pro A	GTC (TAGT	GAGAACATGT	TGGTC	TACGCTTTCT

4IG. 3F

GGCGCCCCGG TACCTCTAGA CCTGGCGATT CAACGTGGTC GGATCATGAC GCTTCCAGAA	09
AACATCGAGC AAGCTCTCAA AGCTGACCTC TTTCGGATCG TACTGAACCC GAACAATCTC	120
GTTATGTCCC GTCGTCTCCG AACAGACATC CTCGTAGCTC GGATTATCGA CGAATCCATG	180
GCTATACCCA ACCTCCGTCT TCGTCACGCC TGGAACCCTC TGGTACGCCA ATTCCGCTCC	240
CCAGAAGCAA CCGGCGCCGA ATTGCGCGAA TTGCTGACCT GGAGACGGAA CATCGTCGTC	300
GGGTCCTTGC GCGATTGCGG CGGAAGCCGG GTCGGGTTGG GGACGAGACC CGAATCCGAG	360
CCTGGTGAAG AGGTTGTTCA TCGGAGATTT ATAGACGGAG ATGGATCGAG CGGTTTTGGG	420
GAAAGGGGAA GTGGGTTTGG CTCTTTTGGA TAGAGAGAGT GCAGCTTTGG AGAGAGACTG	480
GAGAGGTTTA GAGAGACG CGGCGGATAT TACCGGAGGA GAGGCGACGA GAGATAGCAT	540
TATCGAAGGG GAGGGAGAAA GAGTGACGTG GAGAAATAAG AAACCGTTAA GAGTCGGATA	009

FIG. 4A

TTTATCATAT TAAAAGCCCA ATGGGCCTGA ACCCATTTAA ACAAGACAGA TAAATGGGCC	ATGGGCCTGA	ACCCATTTAA	ACAAGACAGA	TAAATGGGCC	099
TAACAGAGT	GTTAACGTTC	GTGTGTTAAG TTAACAGAGT GTTAACGTTC GGTTTCAAAT GCCAACGCCA TAGGAACAAA	GCCAACGCCA	TAGGAACAAA	720
CTCAAGTAA	ACCCCTGCCG	ACAAACGIGT CCTCAAGTAA ACCCCTGCCG TTTACACCTC AATGGCTGCA TGGTGAAGCC	AATGGCTGCA	TGGTGAAGCC	780
CGTAGGAT	GCATGACGAC	ATTAACACGT GGCGTAGGAT GCATGACGAC GCCATTGACA CCTGACTCTC TTCCCTTCTC	CCTGACTCTC	TTCCCTTCTC	840
TAATCAAT	TCAACTACTC	TTCATATATC TCTAATCAAT TCAACTACTC ATTGTCATAG CTATTCGGAA AATACATACA	CTATTCGGAA 1	AATACATACA	006
TTCGATCT	CTCTCAATTC	CATCCTITIC ICTICGAICT CICTCAAITC ACAAGAAGCA AAGICGACGG AICCCIGCAG	AAGTCGACGG 1	ATCCCTGCAG	096
IGACTATT .	TTCATAGTCC	TAAATTACGC CATGACTATT TTCATAGTCC AATAAGGCTG ATGTCGGGAG TCCAGTTTAT	ATGTCGGGAG 1	rccagititat	1020
STTTAGAA	TTTGATCAAT	GAGCAATAAG GTGTTTAGAA TTTGATCAAT GTTTATAATA AAAGGGGGAA GATGATATCA	AAAGGGGAA G		1080
CTTTTTGG (CTTTTGTTAA	CAGTCTTTTG ITCTTTTGG CTTTTGTTAA ATTTGTGTGT ITCTATTTGT AAACCTCCTG	TCTATTTGT A	AACCTCCTG	1140
TCTTTCC (TTTTTAAGT	TATATGTTGT ACTTCTTTCC CTTTTTAAGT GGTATCGTCT ATATGGTAAA ACGTTATGTT	ATATGGTAAA A	CGTTATGTT	1200

FIG. 4B

TGGTCTTTCC	TGGTCTTTCC TTTTCTCTGT TTAGGATAAA AAGACTGCAT GTTTTATCTT TAGTTATATT	TTAGGATAAA	AAGACTGCAT	GTTTTATCTT	TAGTTATATT	1260
ATGTTGAGTA	ATGITGAGTA AATGAACTIT CATAGATCTG GTTCCGTAGA GTAGACTAGC AGCCGAGCTG	CATAGATCTG	GTTCCGTAGA	GTAGACTAGC	AGCCGAGCTG	1320
AGCTGAACTG	AGCTGAACTG AACAGCTGGC AATGTGAACA CTGGATGCAA GATCAGATGT GAAGATCTCT	AATGTGAACA	CTGGATGCAA	GATCAGATGT	GAAGATCTCT	1380
AATATGGTGG	AATATGGTGG TGGGATTGAA CATATCGTGT CTATATTTTT GTTGGCATTA AGCTCTTAAC	CATATCGTGT	CTATATTTT	GTTGGCATTA	AGCTCTTAAC	1440
АТАСАТАТАА	ATAGATATAA CTGATGCAGT CATTGGTTCA TACACATATA TAGTAAGGAA TTACAATGGC	CATTGGTTCA	TACACATATA	TAGTAAGGAA	TTACAATGGC	1500
AACCCAAACT	AACCCAAACT TCAAAAACAG TAGGCCACCT GAATTGCCTT ATCGAATAAG AGTTTGTTTC	TAGGCCACCT	GAATTGCCTT	ATCGAATAAG	AGITITGITITC	1560
CCCCCACTTC	CCCCCACTTC ATGGGATGTA ATACATGGGA TTTGGGAGTT TGAATGAACG TTGAGACATG	ATACATGGGA	TTTGGGAGTT	TGAATGAACG	TTGAGACATG	1620
GCAGAACCTC	GCAGAACCTC TAGAGGTACC GGCGCGC	ටවටවටව				1647

48	96	144	192	240	28 88	336
GTT Val	TTC	AAC Asn	Grr Val	GTT Val	GAT Asp	TGC
ATT Ile	ACG Thr	GTA Val	GCG Ala	GAA Glu	TAT Tyr	GTT :
GAG Glu	CCA Pro	TCC Ser	CAT His	GCC	GAC ASP	ACC (Thr 1
ACC Thr	TCA	CAA Gln	AAC Asn	AGT (TGG (Trp /	TTG 7 Leu 1
TCT Ser	GGT Gly	CTT	ATA Ile	TTT Phe	CTT :	GTC 1 Val 1
CTC	GCC Ala	TTT Phe	CTC Leu	GTG	AAG (Lys]	TTT (Phe 1
CTA Leu	AAC Asn	GAT ASD	TAC Tyr	CTT	AAG Lys]	GTC :
GAT Asp	CCA Pro	CCG Pro	CAC His	GTG	TGG 7	GGT (
CAA Gln	$_{\rm GGT}^{\rm GGT}$	TTA Leu	TAT Tyr	CTT	ATT 1	TTC (Phe (
GAA Glu	TCC Ser	CGT	$_{\rm G1y}^{\rm GGT}$	GTT Val	GAG Glu	TTC :
AGC	CCT	AGA Arg	CTT Leu	CCG	GAA Glu (GGA G
TCT	GAA Glu	CGG Arg	AAA Lys	ATA Ile	GGA (ATC (Ile (
AGG Arg	ATC Ile	GTC Val	GTG Val	ACG	AGC (GTC 1 Val
AGT Ser	GGG	AGA Arg	TAC Tyr	GCG Ala	TTA Leu	ACC (Thr 1
ATG Met	CGT Arg	GTC Val	AAG Lys	TTG	AGT Ser]	GCA A
GAA	AAC Asn	TCG Ser	TTĞ Leu	TAC	GGG 7	ATC (Ile A

FIG. 5A

384	432	480	528	576	624	672	720
ATG TCT CGT CCA CGA TCT GTT TAT CTC ATT GAC TTC GCT 384 Met Ser Arg Pro Arg Ser Val Tyr Leu Ile Asp Phe Ala	CCT TCC GAT GAA CTT AAG GTG ACA AGA GAA GAG TTC ATA 432 Pro Ser Asp Glu Leu Lys Val Thr Arg Glu Glu Phe Ile	AGA AAA TCA GGC AAG TTC GAC GAA GAG ATC CTC GGA TTC 480 Arg Lys Ser Gly Lys Phe Asp Glu Glu Ile Leu Gly Phe	ATC CTT CAA GCC TCA GGA ATA GGC GAT GAA ACG TAC GTC 528 Ile Leu Gln Ala Ser Gly Ile Gly Asp Glu Thr Tyr Val	ATC TCT TCG TCG GAA AAC ACA ACA ACG ATG AAA GAA GGT 576 Ile Ser Ser Ser Glu Asn Thr Thr Thr Met Lys Glu Gly	GCC TCG ATG ATA TTC GGC GCA CTC GAC GAA CTC TTC 624 Ala Ser Met Met Ile Phe Gly Ala Leu Asp Glu Leu Phe	CGT GTC AAA CCG AAA GAC GTA GGT GTC CTC GTG GTT AAC 672 Arg Val Lys Pro Lys Asp Val Gly Val Leu Val Val Asn	TTT AAC CCG ACT CCG TCA CTC TCC GCG ATG GTG ATT AAC 720 Phe Asn Pro Thr Pro Ser Leu Ser Ala Met Val Ile Asn
TTC	AAG Lys	GCT Ala	AGG	Ser	GAA (Glu 2	ACA (Thr A	ATC T Ile F
TAC T Tyr P	TTC A Phe L	CTA G Leu A	AAG A(Lys Aı	AGA TC Arg Se	GAA GA Glu Gl	AAG AC Lys Th	AGT AT Ser Il
GTC Val	TGT	GAT (Asp]	AAG A Lys I	CCA Pro Pro P	CGT G	GAG A Glu L	TGC A Cys S

768	816	864	912	096	1008	1056	1104
ATG Met	CIT	GTT Val	AAC Asn	CGC	ACT Thr	GAA Glu	GAA
GGG G1y	ATG Met	ATG Met	CCT	CGC Arg	CGG	GAA (ATG (Met (
GGA Gly	GAC Asp	GAG Glu	ATA Ile	AAC Asn	GTC	CAG (Glu (CTA 7
CTA	CGT	ACC	GTT Val	TCT Ser	ATT (Ile	TAC (Tyr (GAC (Asp I
AAC Asn	GCT Ala	AGT Ser	ATG Met	CTG	CAC	GTG '	AGA (Arg 1
TAC	CTT Leu	GTG Val	TCA Ser	ATG Met	GAG Glu	AGT	AGC
AGC	GAT ASP	GTT Val	AAG Lys	GTT Val	CTT Leu	AGG Arg	ATA Ile
CTT	GTT Val	GTG Val	GAC Asp	GCC Ala	CGC	TTC	AAA Lys
ATA Ile	GCC Ala	GCG	CGT	TCC Ser	TAC	AGC	TTA
AAC Asn	ATA Ile	TAC	GGA Gly	TGC Cys	AAG Lys	CGT Arg	GGA G1y
GGG G1y	ATC Ile	AGT Ser	GTG Val	GGT Gly	GCT Ala	GAC Asp	AAG
AGA Arg	GGA Gly	AAT Asn	TAC	ATG Met	CAT His	GAC Asp	TTC
ATG Met	GCA Ala	CCG	TGG	AGG Arg	CGC Arg	GCC Ala	GGA G1y
AAG Lys	TCA	AAC Asn	AAT Asn	TTT Phe	TTC	GCT	CAA G1n
TAC	TGC	TCT Ser	TAT Tyr	TTC	GAC Asp	AAG Lys	GAA Glu
CAC His	GGT Gly	CAG Gln	$\texttt{GGG}\\ \texttt{G1y}$	TGC Cys	CGT Arg	CAC His	GAT Asp

	FIG. 5D
1488	GGA AAC ACT TCC AGC AGT GGA ATC TGG TAC GAG CTT GCT TAC ATG GAG Gly Asn Thr Ser Ser Ser Gly Ile Trp Tyr Glu Leu Ala Tyr Met Glu
1440	TTG AGT GAT GAG AAC ATG GAG GCT TCT AAG ATG ACT TTA CAC AGG TTT Leu Ser Asp Glu Asn Met Glu Ala Ser Lys Met Thr Leu His Arg Phe
1392	CAC GCG GCA AGC AAA GCG GTG CTT GAG GAG CTT CAG AAG AAT CTA GGC His Ala Ala Ser Lys Ala Val Leu Glu Glu Leu Gln Lys Asn Leu Gly
1344	AAG CCG TAC ATC CCG GAC TAC AAG CTT GCC TTC GAG CAT TTC TGC TTC Lys Pro Tyr Ile Pro Asp Tyr Lys Leu Ala Phe Glu His Phe Cys Phe
1296	ACT GCG AAA ATC AAC GGA GCC AAG TCG TCA TCC TCC TCT GAT CTA TCC Thr Ala Lys Ile Asn Gly Ala Lys Ser Ser Ser Ser Ser Asp Leu Ser
1248	AGA ACT TTC TCA CCC GCC GCC AAA ACT ACC ACC ACC TCC TCC TCA GCC Arg Thr Phe Ser Pro Ala Ala Lys Thr Thr Thr Thr Ser Ser Ala
1200	GTC CTT CCT TTC TCC GAG CAG CTT CTC TTC TTT GCC GCT TTG ATC CGT Val Leu Pro Phe Ser Glu Gln Leu Leu Phe Phe Ala Ala Leu Ile Arg
1152	GTT GGA GGT GAA GCT CTC AAG ACC AAC ATC ACC ACC TTA GGC CCT CTC Val Gly Gly Glu Ala Leu Lys Thr Asn Ile Thr Thr Leu Gly Pro Leu

1536	1584	1632	1687	1747	1807	1810
GCC AAG GAG AGT GTT CGT AGA GGC GAT AGG GTT TGG CAG ATT GCT TTT Ala Lys Glu Ser Val Arg Arg Gly Asp Arg Val Trp Gln Ile Ala Phe	GGG TCA GGT TTT AAG TGT AAC AGT GTG GTT TGG AAG GCA ATG AGG AAG Gly Ser Gly Phe Lys Cys Asn Ser Val Val Trp Lys Ala Met Arg Lys	GTG AAG AAG CCG GCA AGG AAC AAT CCT TGG GTT GAT TGC ATT AAC CGT Val Lys Lys Pro Ala Arg Asn Asn Pro Trp Val Asp Cys Ile Asn Arg	TAC CCT GTC GCT CTC TGATCATTTA TTTTTAAAAT TATTATTTCT TCTTAATTAA	ATCATCTATG ATCTCTTTC CTTGTTGTTG GATGATAGAC GTTTGTTTGC TGGTCATTCG	TATCTTAAGA CTTCTATAAG AATGGATGGT TCAAGTCCAA AAAAAAAA AAAAAAAAA 1807	AAA

FIG. SE

51	66	147	195	243	291	339
GTCGACAAA ATG ACG TCC ATT AAC GTA AAG CTC CTT TAC CAT TAC GTC ATA Met Thr Ser Ile Asn Val Lys Leu Leu Tyr His Tyr Val Ile	ACC AAC CTT TTC AAC CTT TGT TTC TTT CCA TTA ACG GCG ATC GTC GCC Thr Asn Leu Phe Asn Leu Cys Phe Phe Pro Leu Thr Ala Ile Val Ala	GGA AAA GCC TAT CGG CTT ACC ATA GAC GAT CTT CAC CAC TTA TAC TAT Gly Lys Ala Tyr Arg Leu Thr Ile Asp Asp Leu His His Leu Tyr Tyr	TCC TAT CTC CAA CAC AAC CTC ATA ACC ATT GCT CCA CTC TTT GCC TTC Ser Tyr Leu Gln His Asn Leu Ile Thr Ile Ala Pro Leu Phe Ala Phe	ACC GTT TTC GGT TCG GTT CTC TAC ATC GCA ACC CGG CCC AAA CCG GTT Thr Val Phe Gly Ser Val Leu Tyr Ile Ala Thr Arg Pro Lys Pro Val	TAC CTC GTT GAG TAC TCA TGC TAC CTT CCA CCA ACG CAT TGT AGA TCA Tyr Leu Val Glu Tyr Ser Cys Tyr Leu Pro Pro Thr His Cys Arg Ser	AGT ATC TCC AAG GTC ATG GAT ATC TTT TAC CAA GTA AGA AAA GCT GAT Ser Ile Ser Lys Val Met Asp Ile Phe Tyr Gln Val Arg Lys Ala Asp

FIG. 6A

387	435	483	531	579	627	675	723
TTG	CCC	CGT Arg	AAG Lys	TCA	ACT Thr	GGT	CAT His
TTC	666	GCG Ala	TTC	AAC	AAC Asn	ATG Met	TTG
GAC Asp	CAC His	GCG	CTA	GTG Val	GTT Val	GGC Gly	TTG
CTT	ACC	GCG Ala	AAT Asn	GTG Val	GTC Val	$_{\rm G1y}^{\rm GGT}$	GAC
TGG	GAA Glu	TTT Phe	GAA	CTT	ATG	CTT	AAG (Lys
TCC	GAT Asp	ACT Thr	CTA	ATA Ile	GCG	AAC	GCA Ala
TCG	GGC Gly	AAG Lys	GCG Ala	$_{\rm GLY}^{\rm GGT}$	TCC Ser	TTT	CTA
GAC Asp	CTA	CGG Arg	GGT G1y	ATA Ile	CTC	AGC Ser	GAT
GAT Asp	GGT G1y	CCC Pro	ATT Ile	GAT Asp	TCG Ser	AGA Arg	ATT Ile
TGC Cys	TCA Ser	CCT	ATC Ile	AAA Lys	CCT	GTA Val	GCC
ACG Thr	CGT Arg	GTC Val	GTT Val	CCT Pro	ACT Thr	AAC Asn	ATA Ile
GGC Gly	GAA Glu	CAG Gln	CAA Gln	AAC Asn	CCA	AGC Ser	GTT Val
AAC Asn	CAA Gln	CTT Leu	GAG Glu	GTT Val	AAT Asn	CGA Arg	GGC G1y
CGG Arg	ATT Ile	CTG	ACG Thr	AAT Asn	TTT Phe	CTC	GCC Ala
TCT	AAG Lys	666	GAG Glu	ACC	ATG Met	AAG Lys	AGT Ser
CCT	AGG Arg	GAG Glu	GAA Glu	AAC Asn	AGC Ser	TTC Phe	TGT Cys

TGC 819 Cys AGA 867 Arg			CAT 915 His	GTT 963 Val	GTT 1011 Val	ATT 1059 Ile	AAG 1107 Lys	
AAT ASN CCT Pro Pro Thr I				GAC (Asp 1	GAT C	TTG A	GGC A Gly L	
TCA Ser AAG Lys CGA	• •			GGA (ACC (Thr 1	CCG 1 Pro I	ATG G Met G	
GTT Val AAC Asn GTT	AAC Asn GTT Val	GTT Val		CAA Gln	ATA Ile	GGT Gly	TTC /	
ATG Met TCC Ser ACG				CAA Gln	GAC Asp	CTG	ACC	
ATG CTC Leu CAC				GTG Val	AAG Lys	ACG	GTT Val	
TCC Ser TTG Leu GTT				TGC	TCC Ser	GCA Ala	TTC	7
AGG Arg ATT Ile CTA Leu		CTA Leu		CGT Arg	TTG	ATA Ile	TTT Phe	ر ازا
AAT Asn GCT Ala GAG Glu		GAG Glu		TTT Phe	AGT Ser	AAC Asn	CIT	-
			Tyr	TCT Ser	GTG Val	AAA Lys	CIT	
GGT GLY GGG GLY			AAG Lys	AAG Lys	GGA G1y	AAG	AAA Lys	
GCT Ala GGT G1y	GGT G1y	٠	TCC	GAC Asp	ACC Thr	GTT Val	GAG Glu	
TAC Tyr GTT Val			CGG Arg	GAC Asp	AAA Lys	ACG Thr	AGC	
ATT Ile		CGT	AGA Arg	GCT Ala	GGC Gly	CGA Arg	TTA	
	Asn	TTC	CGT Arg	GGA Gly	AAC Asn	$_{\rm GIY}^{\rm GGT}$	CCG	
	TAT TYY	TTG	GAT Asp	ACC Thr	GAG Glu	GCT Ala	CTT Leu	

1155	1203	1251	1299	1347	1395	1442
AAG Lys	ATT Ile	GCA Ala	ATA Ile	GGT Gly	AGT Ser	
TTC	GTG Val	GAG Glu	TCA	AAA Lys	AAC Asn	ATCC
GAC	GCC Ala	GTA Val	AGC	AAG Lys	TGT	TAGGATCC
CCG Pro	AAA Lys	GAT Asp	TCT Ser	ATG Met	AAG Lys	AAA '
GTC Val	GGC Gly	ATC Ile	TCA	AGG	TTT	TCC
TAC	${\tt GGA}$	CCG Pro	ACT	GGA Gly	GGC G1y	GCT
TAT Tyr	GCC Ala	GCA Ala	AAC Asn	AAA Lys	TCA Ser	AAA
CAT His	CAT	CTA Leu	GGA Gly	GCA Ala	666 G1y	GTC Val
AAA Lys	ATA Ile	GGC G1y	TTT Phe	GAA Glu	TTA Leu	AAT Asn
ATC Ile	TGT Cys	CTA Leu	AGA Arg	ATA Ile	GCT	AAC Asn
AAA Lys	TTT Phe	AAC Asn	CAT His	TAC	ATT	CTA Leu
GAC	CAT	AAG Lys	TTA Leu	GCA Ala	CAG Gln	GCT Ala
AAA Lys	GAC Asp	GAG Glu	ACG Thr	TTG	TGG	GTG Val
TTC	ATC Ile	CTA	TCA	GAG	GTT Val	TGG
CTT	GCT	GTG Val	AGA Arg	TAT Ty <i>r</i>	AAA Lys	GTT Val
AAA Lys	CIT	GAT Asp	TCA	TGG	AAT Asn	GCA Ala

FIG. 6L

51	66	147	195	243	291	339
GTCGACAAA ATG ACG TCC ATT AAC GTA AAG CTC CTT TAC CAT TAC GTC ATA Met Thr Ser Ile Asn Val Lys Leu Leu Tyr His Tyr Val Ile	CTT TTC AAC CTT TGC TTC TTT CCG TTA ACG GCG ATC GTC GCC Leu Phe Asn Leu Cys Phe Phe Pro Leu Thr Ala 11e Val Ala	GCC TAT CGG CTT ACC ATA GAC GAT CTT CAC CAC TTA TAC TAT Ala Tyr Arg Leu Thr Ile Asp Asp Leu His His Leu Tyr Tyr	CTC CAA CAC AAC CTC ATA ACC ATC GCT CCA CTC TTT GCC TTC Leu Gln His Asn Leu Ile Thr Ile Ala Pro Leu Phe Ala Phe	TTC GGT TCG GTT CTC TAC ATC GCA ACC CGG CCC AAA CCG GTT Phe Gly Ser Val Leu Tyr Ile Ala Thr Arg Pro Lys Pro Val	GTT GAG TAC TCA TGC TAC CTT CCA CCA ACG CAT TGT AGA TCA Val Glu Tyr Ser Cys Tyr Leu Pro Pro Thr His Cys Arg Ser	TCC AAG GTC ATG GAT ATC TTT TAT CAA GTA AGA AAA GCT GAT Ser Lys Val Met Asp Ile Phe Tyr Gln Val Arg Lys Ala Asp
CGAC	AAC Asn	A AAA 7 Lys	TAT	GTT Val	CTC	ATC Ile
Ę.	ACC Thr	GGA Gly	TCC Ser	ACC Thr	TAC Tyr	AGT Ser

FIG. 7A

387	435	483	531	579	627	675	723
TTG	CCC	CGT Arg	AAG Lys	TCA Ser	ACT Thr	GGT Gly	CAT His
TTC	666	GCG Ala	TTC Phe	AAC Asn	AAC Asn	ATG Met	TTG
GAC ASP	CAC His	GCG Ala	CTA Leu	GTG Val	GTT Val	GGC G1y	TTG
CTT	ACT Thr	GCG Ala	AAT Asn	GTG Val	GTC Val	GGT Gly	GAC
TGG	GAA Glu	TTT Phe	GAA Glu	CTT	ATG Met	CTT (Leu (AAG (Lys 2
TCG Ser	GAT Asp	ACT Thr	CTA Leu	ATA Ile	GCG Ala	AAC (Asn]	GCA A
TCG	GGC Gly	AAG Lys	GCG Ala	GGT Gly	TCC	TTT . Phe .	CTA (Leu
GAC ASP	CTA	CGG Arg	GGT Gly	ATA Ile	CTC	AGC Ser	GAT
GAT ASP	GGT Gly	CCC Pro	ATT Ile	GAT Asp	TCG Ser	AGA Arg	ATT Ile
TGC Cys	TCA Ser	CCT	ATC Ile	AAA Lys	CCA Pro	GTA Val	GCC
ACG Thr	CGT Arg	GTC Val	GTT Val	CCT Pro	ACT Thr	AAC Asn	ATA Ile
GGC Gly	GAA Glu	CAG Gln	CAA Gln	AAC Asn	CCA Pro	AGC Ser	GTT Val
AAC Asn	CAA Gln	CTT	GAG Glu	GTT Val	AAT Asn	CGA Arg	GGC G1y
CGG Arg	ATT Ile	CTG	ACG Thr	AAC Asn	TTT Phe	CTC	GCC Ala
TCT Ser	AAG Lys	666 G1y	GAG Glu	ACC Thr	ATG Met	AAG Lys	AGT Ser
CCT	AGG Arg	GAG Glu	GAA Glu	AAC Asn	AGC Ser	TTC Phe	TGT Cys

771	819	867	915	963	1011	1059	1107
ACT Thr	TGC	GGA Gly	CAT His	GAT Asp	GTT Val	ATT Ile	AAG Lys
ATC Ile	AAT Asn	CCT Pro	ACG	GAC	GAT Asp	TTG 7	GGC 7 Gly 1
AAC Asn	TCA Ser	AAG Lys	CGA	GGA Gly	ACC	CCG Pro 1	ATG (Met (
GAG Glu	GTT Val	AAC Asn	GTT Val	CAA	ATA Ile	GGT (TTC A
ACA Thr	ATG Met	TCC	ACG	CAA Gln	GAC ASP	TTG (Leu (ACC 1 Thr E
AGC	ATG Met	CIC	CAC His	GTG Val	AAG	ACG Thr 1	GTT 1 Val
GTG Val	TCC	TTG	GTT Val	TGC Cys	TCC	GCA Ala	TTC (Phe 1
GTG Val	AGG Arg	ATT Ile	CTA	CGT Arg	TTG	ATA Ile	TTT P
CTT Leu	AAT Asn	GCT Ala	GAG Glu	TTT Phe	AGT Ser	AAC	CTT
GCT Ala	GAT Asp	GCC Ala	TAC Tyr	TCT	GTG Val	AAA Lys	CTT
TAT Tyr	$_{\rm GGT}$	GGG G1Y	AAG Lys	AAG Lys	GGA Gly	AAG Lys	AAA
ACG	GCT Ala	GGT Gly	TCC	GAC Asp	ATC Ile	GTT Val	GAG
AAT Asn	$ ext{TAC}$	GTT Val	CGG Arg	GAC Asp	AAA Lys	ACG Thr	AGC
AAA Lys	ATT Ile	CGT Arg	AGA Arg	GCT Ala	GGC G1y	CGA Arg	TTA
CAT His	AAC Asn	TTC	CGT Arg	GGA G1y	AAC Asn	GGT Gly	CCG Pro
GTC Val	$ extsf{TAT} extsf{T} extsf{T} extsf{Y} extsf{z}$	TTG	GAT Asp	ACC	GAG Glu	GCT Ala	CIT

1155	1203	1251	1299	1347	1395	1442
CTT TTC AAA GAT AAA ATC AAA CAT TAC TAC GTC CCG GAT TTC AAA Leu Phe Lys Asp Lys ile Lys His Tyr Tyr Val Pro Asp Phe Lys	GCT ATT GAC CAT TTT TGT ATA CAT GCC GGA GGC AGA GCC GTG ATT Ala Ile Asp His Phe Cys Ile His Ala Gly Gly Arg Ala Val Ile	GTG CTA GAG AAC CTA GCC CTA GCA CCG ATC GAT GTA GAG GCA Val Leu Glu Lys Asn Leu Ala Leu Ala Pro Ile Asp Val Glu Ala	AGA TCA ACG TTA CAT AGA TTT GGA AAC ACT TCA TCT AGC TCA ATA Arg Ser Thr Leu His Arg Phe Gly Asn Thr Ser Ser Ser Ile	TAT GAG TTG GCA TAC ATA GAA GCA AAA GGA AGG ATG AAG AAA GGT Tyr Glu Leu Ala Tyr Ile Glu Ala Lys Gly Arg Met Lys Lys Gly	AAA GTT TGG CAG ATT GCT TTA GGG TCA GGC TTT AAG TGT AAC AGT Lys Val Trp Gln Ile Ala Leu Gly Ser Gly Phe Lys Cys Asn Ser	GTT TGG GTG GCT CTA AAC AAT GTC AAA GCT TCC AAA TAGGATCC Val Tro Val Ala Leu Asn Asn Val Lvs Ala Ser Lvs
AAA Lys	CTT	GAT ASD	TCA Ser	TGG Trp	AAT Asn	GCA

FIG. 7D

					•	
48	96	144	192	240	288	336
ATC Ile	TTC Phe	TTC	GCT Ala	TGT Cys	GAC Asp	CAA Gln
TAC	TCT Ser	CAT His	ACC Thr	TCG	ATG Met	TTC (Phe (
CTC	TCT	TTC	TCC	TTC	TTC	GCT A
GCT Ala	CTC	CGT Arg	CTC	GAC Asp	ACA Thr	TTA Leu
AAC Asn	AAC Asn	CTC Leu	TCT	CTC	GAA	AAC '
TCC	GCT Ala	ACA Thr	ATC Ile	CTC	CGT Arg	GAC
ATC	ATC Ile	AAC Asn	TTG	TTC	ACT	GAA Glu
CTA	ACA	TAC	CTC	GTC Val	TGC	ACA Thr
TAC	GCA Ala	CTC	GCA Ala	CGT Arg	ATC Ile	TTC
CAT	GCC Ala	CTC	ACC	CGC	CTG	ATC Ile
TAT Tyr	CTC	TCT Ser	GCC Ala	CCT Pro	TCA Ser	GGC Gly
GTG	CIC	CTC	CTC	CGT Årg	CCT	GTA Val
CTA	CCT	GAC	ACA Thr	ACC	GAC Asp	CGT Arg
AAA Lys	CTT	AAC Asn	GCC Ala	ACC Thr	CCA	CAA Gln
CTT	CTC	ATC Ile	TCC	TTC	AAA Lys	TCT Ser
AAG Lys	CTC	ACC	CTC	TAC	$\mathtt{TAC}\\ \mathtt{TYr}$	AGA Arg

384	432	480	528	576	623
CCT Pro	AGA Arg	GAG Glu	TGT Cys	AAG Lys	GG G1<
TTC	GCG Ala	CTT Leu	AAT Asn	AAT Asn	ATG
TAC	GAA Glu	GTT Val	GTG Val	GTG Val	GGG G1v
ACT Thr	GAA Glu	GCG Ala	GTG Val	ATT Ile	GGC (
AAA Lys	ATG Met	GAC Asp	CTT Leu	ATG	TTC (Phe (
CAG Gln	TGT Cys	ATT Ile	ATC Ile	GCT	AAT '
GGT Gly	CCT	GCT Ala	$_{\rm GLY}^{\rm GGA}$	TCT	TAT
CTA	AAT Asn	GGA Gly	ATT Ile	CTT	AGC
GGT G1y	CCT	TTC Phe	gat Asp	TCA Ser	TTG
TCC	CCT	ATG Met	AAA Lys	CCG Pro	ATT
AGA Arg	GTT Val	GTT Val	CCT	ACA Thr	AAC
GAA Glu	CGT Arg	ACA Thr	AAA Lys	CCA Pro	GGC
CIC	CTT Leu	GAA Glu	GTG Val	AAT Asn	AGA Arg
ATC Ile	CTT Leu	GCA Ala	$_{\rm GGT}^{\rm GGT}$	TTT	CTT
AAG Lys	GCT Ala	GAG Glu	ACC	TTG	AAG Lys
CAA Gln	GAA Glu	AAA Lys	AAG Lys	AGC	TAT

48	96	144	192	240	288	336	384
CTC	TTA Leu	TTC	CGA Arg	TCG Ser	ATT Ile	ATT Ile	TCT Ser
AAA Lys	TTG	ATC Ile	TCT Ser	CCT	TTG /	TTG 7	CAC T His S
TTT Phe	TCA	TTC Phe	ATG Met	CCG	AGT '	ATC : Ile I	ATT C Ile H
TTT Phe	GTC Val	$_{\rm GGA}$	TTC	CTC	TCT Ser	AAG 1 Lys	TCT A
CAC His	AAT Asn	ACC	TTC	TAC	AAC A	AGG 1 Arg I	GAT 1 ASP S
ACT	ATG Met	TCC Ser	GTC Val	TGC Cys	AAC	CAG Z	CCG (Pro Pro P
ATC Ile	TTC	AAT Asn	ATT Ile	TCT Ser	ATG Met	TTC	TTA (Leu]
CTG	TTG Leu	$ ext{TAC}$	TCC Ser	\mathtt{TAC}	TTC	GAG	TAT Tyr] FIG.
TAT	GTT Val	TAT Tyr	GGA Gly	GAT Asp	AAA Lys	CTT	ACT ' Thr '
CAC His	GCT Ala	CTC	GTC Val	CTA Leu	CAG Gln	TCT	GAG Glu
TAC	ATG Met	CAG Gln	ATT Ile	CTT Leu	$\mathtt{TAC}\\ \mathtt{TYr}$	ACT	GAA Glu
GGC G1y	CTA	CTT	GCC Ala	TAC	AGC Ser	GAA Glu	GGT G1y
TTA	CCT	CAT His	CTC	ATC Ile	GTT Val	AGC Ser	CTC
AAG Lys	CTC	AAC Asn	ACT Thr	TCC	AAA Lys	TTC	GGT Gly
CIT Leu	Phe	CTA	ATC Ile	AGA Arg	CAA Gln	GAT ASp.	TCT (
AAG Lys	ATG Met	AGC Ser	GTC Val	CCT	AGT Ser	CAA Gln	CGC

432	480	528	576	209
CAG	AAT	CCC	GGA	
Gln	Asn	Pro	G1y	
GAG	ATC	AAC	AGA	
Glu	Ile	Asn	Arg	
GĆG Ala	AAA Lys	TTT Phe	CTT	
GAA	ACA	TTG	AAG	
Glu	Thr	Leu	Lys	
GAA Glu	AAT Asn	AGT Ser	TAT	
CGT	GAG	TGT	AAG	ტ
Arg	Glu	Cys	Lys	
GCG	TTC	AAT	AAC	ATG
Ala	Phe	Asn	Asn	
GCA	CTT	GTG	GTT	GGC
Ala	Leu	Val	Val	
GCT	AAT	GTT	ATT	GGC
Ala	Asn	Val	Ile	G1v
ATG	GAC	CTT	ATG	CTC
Met	Asp	Leu	Met	
ACT	CTC	GTT	GCC	AAT
	Leu	Val	Ala	Asn
CCT	GCA Ala	$_{\rm GLY}^{\rm GGT}$	TCC	TTT Phe
CGT	$_{\rm G1Y}^{\rm GGT}$	ATT Ile	TTA Leu	AGC Ser
- CCG	TTC	GAG	TCT	AAG
Pro		Glu	Ser	Lys
CCT	ATC Ile	AGG Arg	CCT	ATT Ile
ATC Ile	GTA Val	CCT	ACG	AAC Asn

48	96	144	192	240	288	336
AAA CTG GGG TAC CAC TAC CTC ATT ACT CAT CTC TTC AAG CTC 48 Lys Leu Gly Tyr His Tyr Leu Ile Thr His Leu Phe Lys Leu	GTT CCA TTA ATG GCG GTT TTA GTC ACA GAG ATC TCC CGA TTA 96 Val Pro Leu Met Ala Val Leu Val Thr Glu Ile Ser Arg Leu	GAC GAT CTT TAC CAG ATT TGC CTT CAT CTC CAA TAC AAT CTC 144 ASP ASP Leu Tyr Gln Ile Cys Leu His Leu Gln Tyr Asn Leu	TTC ATC TTT CTC TCT GCT TTA GCT ATC TTT GGC TCC ACC GTT 192 Phe Ile Phe Leu Ser Ala Leu Ala Ile Phe Gly Ser Thr Val	ATG AGT CGT CCC AGA TCT GTT TAT CTC GTT GAT TAC TCT TGT 240 Met Ser Arg Pro Arg Ser Val Tyr Leu Val Asp Tyr Ser Cys	CCT CCG GAG AGT CTT CAG GTT AAG TAT CAG AAG TTT ATG GAT 288 Pro Pro Glu Ser Leu Gln Val Lys Tyr Gln Lys Phe Met Asp	AAG TTG ATT GAA GAT TTC AAT GAG TCA TCT TTA GAG TTT CAG 336 Lys Leu Ile Glu Asp Phe Asn Glu Ser Ser Leu Glu Phe Gln
AAG CTT Lys Leu	TGT TTG Cys Leu	ACA ACA Thr Thr	GTT GCT Val Ala	TAC ATC I	TAT CTT (Tyr Leu	CAT TCT A

384	432	480	528	576	622
CCT	CGT	GAG Glu	TGT Cys	AAG Lys	ტ
CIC	GCT Ala	TTC	AAT Asn	AAC Asn	ATT Ile
TAT Tyr	GCG Ala	CTT Leu	GTG Val	GTT Val	66C 61y
ACT Thr	ATG Met	AAG Lys	GTT Val	ATT Ile	GGG G1y
GAG Glu	ATG Met	GAT	TTG Leu	ATG Met	CTG
GAA	ACG	CTT	GTG Val	GCT	AAC Asn
GGA Gly	CCT Pro	GCT Ala	$_{\rm GLY}^{\rm GGT}$	TCA	TTT Phe
TTA	AGG Arg	$_{\rm GGT}$	ATT Ile	TTG Leu	AGT Ser
GGT G1y	CCG	TTT Phe	GAT	TCG Ser	AAG Lys
TCT Ser	CCT	ATG Met	AGG Arg	CCT	GTT Val
CGT	ATC Ile	GTA Val	CCT	ACA Thr	AAT Asn
GAA	TGT Cys	CAG Gln	AAC	CCT	GGG G1y
· CTT Leu	CAT	GAG	ATT Ile	AAT Asn	AGA Arg
ATT Ile	TTA Leu	GCT	AAG Lys	TTT Phe	CTT Leu
AAG	GCT Ala	GAA Glu	ACC	TTG	AAG Lys
AGG Arg	GAA Glu	GAG Glu	AAT Asn	AGC	TAT TYT

FIG. 10]

48	96	144	192	240	288	336
CTC	TTA Leu	AAT Asn	ACA Thr	TCA Ser	ATG Met	TTC Phe
AAG Lys	CGG Arg	TTC	TTC Phe	TAC Tyr	TTC	GAG (
TTT Phe	TCC Ser	CAG Gln	GGA Gly	GAC '	ACA T	CTT (
CTT Leu	GTC Val	CTC Leu (TTC (Phe (CTC (Leu)	CAG A	TCG C Ser L
CAC (His]	AAT (Asn 1	CAG (Gln I	ATT 1 Ile E	CTC C Leu I	TAC C Tyr G	TCG T Ser S
TCT (ACG 1 Thr 1	CTC C	TCC A Ser I	TAC C	AGC T	GAG T Glu S
ATT I	TTC 1 Phe 1	TCT (Ser I	GTC 1 Val S	GTT T Val T	GTT A Val S	GAC G. Asp G
CTG 7	CTG 1 Leu 1	CTC 7 Leu 9	ACC G Thr V	TCC G Ser V	AAA G Lys V	TTC G
TAC (Tyr I	GTT C Val 1	GAT C ASP L	ATT A Ile T	AGA T Arg S	CTC A	
CAC 1 His 1	GCG G Ala V	CTC G Leu A	TTC A	CCT A		A GAT u Asp
TAT C Tyr H	ATG G Met A				G AAT r Asn	r GAA e Glu
		TGT Cys	: TTC	CGA	TCG	ATT Ile
TGG Trp	TTA Leu	CTC	ATC Ile	TCC Ser	CCG Pro	CTG
TTA Leu	CCT	CAG Gln	TTC Phe	ATG Met	CCG Pro	AAA Lys
AAG Lys	GTT Val	AAC Asn	GGA Gly	TTC	CTC	TCT Z
CTT	TTG	CTA	GTC	ATC Ile	TAC (Tyr 1	CAT :
AAG Lys	TTG	AGC	CTC	GTT 7	TGT C	AAT (Asn F

416. 114

4	2	1	59
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384	432	480	528	576	625
CTC	GCG Ala	TTC	AAC Asn	AAC Asn	ATG G Met>
TAC (Tyr I	GCG C	CTC 1 Leu F	GTG A Val A	GTG A Val A	GGC A Glv M
ACT Thr	GCG Ala	AAT	GTG (Val	ATT (Ile 1	GGT (
GAG	ATG Met	GAC	GTG Val	ATG Met	CTC Leu
GAA	ACT Thr	CTC	GTT Val	GCC Ala	AAT Asn
3 GGC 1 G1Y	CCG Pro	GCA Ala	GGT	TCC	TTT Phe
r CTC 7 Leu	3 CGT 5 Arg	GGT G1y	ATT 1 Ile	TTA Leu	AGC
c GGT r Gly	3 CCG 5 Pro	TTC Phe	s GAG y Glu	r TCT Ser	AAG Lys
A TCC g Ser	c ccG e Pro	A ATC 1 Ile	r AGG o Arg	3 CCT r Pro	C GTG
G CGA	C ATC	G GTA n Val	C CCT Pro	G ACG o Thr	A AAC V Asn
CTG AAG Leu Lys	CAC TGC His Cys	GAG CAG Glu Gln	ATC GAC Ile Asp	AAC CCG Asn Pro	ia gga og gly
ATC C Ile L	ATC CZ Ile H	TCG GA Ser GJ	AAA A1 Lys I1	TTT A? Phe As	CTT AGA Leu Arg
AAG A Lys I	TCT A Ser I	GAA T	ACC A Thr L	TTG T	AAG C' Lys L
CGG A	GAA 7 Glu 9	GAG (Glu (AAT A Asn 1	AGC 1 Ser I	TAT A
CAG Gln	CCG	CGT	GAG	TGC	AAG '

FIG. 11B

26	104	152	200	248	296	344
GITCAITGAI ITGIITGAGA CICIGITGCA GAAAICICCA C AIG GAI GAI GAA ICC Met Asp Asp Glu Ser	GTT AAT GGA GGA TCC GTA CAG ATC CGG ACC CGA AAG TAC GTC AAG CTG Val Asn Gly Gly Ser Val Gln Ile Arg Thr Arg Lys Tyr Val Lys Leu	GGT TAT CAC TAC CTG ATT TCT CAC CTT TTT AAG CTC TTG TTG GTT CCT Gly Tyr His Tyr Leu Ile Ser His Leu Phe Lys Leu Leu Leu Val Pro	TTA ATG GCG GTT CTG TTC ACG AAT GTC TCC CGG TTA AGC CTA AAC CAG Leu Met Ala Val Leu Phe Thr Asn Val Ser Arg Leu Ser Leu Asn Gln	CTC TGT CTC GAT CTC TCT CTC CAG CTC CAG TTC AAT CTC GTC GGA TTC Leu Cys Leu Asp Leu Ser Leu Gln Leu Gln Phe Asn Leu Val Gly Phe	ATC TTC TTC ATT ACC GCC TCC ATT TTC GGA TTC ACA GTT ATC TTC ATG Ile Phe Phe Ile Thr Ala Ser Ile Phe Gly Phe Thr Val Ile Phe Met	TCC CGA CCT AGA TCC GTT TAC CTC CTC GAC TAC TCA TGT TAC CTC CCG Ser Arg Pro Arg Ser Val Tyr Leu Leu Asp Tyr Ser·Cys Tyr Leu Pro

FIG. 12A

u	11	1	EO	
ч	ч	1	59	

392	440	488	536	584	632	089
AAA Lys	ATC Ile	ATC Ile	TCG	AAA Lys	TTT Phe	CTT Leu
TCT Ser	AAG Lys	TCT	GAA Glu	ACC Thr	TTG Leu	AAG (Lys)
CAT His	CGG Arg	GAA Glu	GAG Glu	AAT Asn	AGC '	TAT 1 Tyr 1
AAT Asn	CAG	CCG	CGT Arg	GAG Glu	TGC Z	AAG 1 Lys 1
ATG Met	TTC Phe	CTC	GCG Ala	TTC Phe	AAC Asn	AAC
TTC Phe	GAG Glu	\mathtt{TAC}	GCG Ala	CTC	GTG	GTG Z
ACA Thr	CTT Leu	ACT	GCG Ala	AAT Asn	GTG Val	ATT (Ile
CAG Gln	TCG Ser	GAG Glu	ATG Met	GAC Asp	GTG Val	ATG Met
TAC	TCG Ser	GAA Glu	ACT Thr	CTC	GTT (Val	GCC ;
AGC Ser	GAG Glu	66C G1y	CCG Pro	GCA	GGT (TCC (
GTT Val	GAC Asp	CTC	CGT Arg	GGT G	ATT (Ile (TTA :
AAA Lys	TTC	$_{\rm GLY}^{\rm GGT}$	CCG Pro	TTC	GAG Glu	TCT Ser 1
CTC	GAT Asp	TCC Ser	CCG Pro	ATC Ile	AGG Arg	CCT
AAT Asn	GAA Glu	CGA Arg	ATC Ile	GTA Val	CCT.	ACG (Thr)
GCG Ala	ATT Ile	AAG Lys	TGC Cys	cAG Gln	GAC (ASp	CCG 7
NCG XXX	CTG	CTG	CAC	GAG (ATC (Ile /	AAC (Asn I

. 12B

			45/59			
728	776	824	872	920	896	1016
GTG AAG AGC TTT AAC CTC GGA GGA ATG GGA TGT AGG GCT	GCC GTT GAT CTC GCT AAT GAC ATT TTA CAG CTC CAT AGA	GCT CTT GTG GTT AGC ACA GAG AAC ATC ACT CAG AAT TGG	GGT AAC AAC AAA GCA ATG TTG ATT CCT AAT TGC TTG TTT AGG	TCC GCG GTT CTG CTT TCG AAC AAG CCT CGT GAT CGA AAA	TAT AAA CTT GTT CAC ACG GTA CGG ACT CAT AAA GGA TCT	sca ttc aac tgt gtg tac caa gaa caa gac gag gac ttg
Val Lys Ser Phe Asn Leu Gly Gly Met Gly Cys Arg Ala	Ala Val Asp Leu Ala Asn Asp Ile Leu Gln Leu His Arg	Ala Leu Val Val Ser Thr Glu Asn Ile Thr Gln Asn Trp	Gly Asn Asn Lys Ala Met Leu Ile Pro Asn Cys Leu Phe Arg	Ser Ala Val Leu Leu Ser Asn Lys Pro Arg Asp Arg Lys	Tyr Lys Leu Val His Thr Val Arg Thr His Lys Gly Ser	ala Phe Asn Cys Val Tyr Gln Glu Gln Asp Glu Asp Leu
AGA GGA AAC	GGT GTC ATC	AAC ACA TTA	TAC TTT GGT	GTT GGT GGA	CGA TCC AAG 1	GAT GAG AAA (
Arg Gly Asn	Gly Val Ile	Asn Thr Leu	TYr Phe Gly	Val Gly Gly	Arg Ser Lys 1	Asp Glu Lys A

			46/59			
1064	1112	1160	1208	1256	1304	1352
CTA ATG TCT ATA GCT GGA GAA	GGT CCT CTG GTT CTT CCA ATA	TTT GTT GCA AAG AGA TTG TTC	TAC ATA CCG GAT TTC AAG CTT	GGA GGT AGA GCC GTG ATC GAT	CCA AAA CAT GTG GAG GCT TCT	ACT TCA TCG AGC TCT ATT TGG
Leu Met Ser Ile Ala Gly Glu	Gly Pro Leu Val Leu Pro Ile	Phe Val Ala Lys Arg Leu Phe	Tyr Ile Pro Asp Phe Lys Leu	Gly Gly Arg Ala Val Ile Asp	Pro Lys His Val Glu Ala Ser	Thr Ser Ser Ser Ile Trp
ACC GGA GTT TCT TTG TCT AAA GAC	CTA AAG ACA AAT ATC ACC ACT TTG	GAG CAG ATT CTG TTC ATT GCG ACT	GCC AAG AAG AAG AAG AAG AAG CCT	TTT GAT CAT TTC TGT ATT CAC GCA	CTA GAG AAG AGT TTA AAG CTA TTG	ATG ACA TTG CAT AGA TTT GGA AAC
Thr Gly Val Ser Leu Ser Lys Asp	Leu Lys Thr Asn Ile Thr Thr Leu	Glu Gln Ile Leu Phe Ile Ala Thr	Ala Lys Lys Lys Lys Lys Pro	Phe Asp His Phe Cys Ile His Ala	Leu Glu Lys Ser Leu Lys Leu Leu	Met Thr Leu His Arg Phe Gly Asn
aaa	GCT	AGC	AGT	GCC	GAA	AGA
Lys	Ala	Ser		Ala	Glu	Arg

TAT GAA TTA GCT TAC ACA GAA GCT AAA GGA AGA ATG AGA AAA GGG AAT Tyr Glu Leu Ala Tyr Thr Glu Ala Lys Gly Arg Met Arg Lys Gly Asn	1400
TGG CAG ATT GCT TTT GGA AGC GGC TTT AAG TGT AAC AGC GCG ITP GIn Ile Ala Phe Gly Ser Gly Phe Lys Cys Asn Ser Ala	1448
GTG GCT CTT CGT GAT GTC GAG CCC TCG GTT AAC AAT CCT TGG Val Ala Leu Arg Asp Val Glu Pro Ser Val Asn Asn Pro Trp	1496
CAT TGC ATC CAT AGA TAT CCG GTT AAG ATC GAT CTC TGATTTCAGC His Cys Ile His Arg Tyr Pro Val Lys Ile Asp Leu	1545
TTAACCGGTA AAATTGGTCT GTACATATAT TTACCACTGA GTAAAGACAT CAGTTAATGA 1605	605
TTTGTTGTTA CTCAATTGGG CTAAGTGTAT TATTATATGT GTTGTATATA ATAAAGGTAG 1665	665
AACGTAAATT TACTAAGAAA AAAAAAAA AAAAAAAAA	1704

FIG. 12E

47	95	143	191	239	287	335
NATG ACG TCT GTG AAC GTA AAA CTC CTT TAC CAT TAC GTC ATA ACC Met Thr Ser Val Asn Val Lys Leu Leu Tyr His Tyr Val Ile Thr	C TTT TTC AAC CTC TGT TTC TTC CCA CTG ACG GGG ATC CTC GCC GGA n Phe Phe Asn Leu Cys Phe Phe Pro Leu Thr Gly Ile Leu Ala Gly	GGC TCT CGT CTT ACC ACA AAC GAT CTC CAC CAC TTC TAT TCA TAT Gly Ser Arg Leu Thr Asn Asp Leu His His Phe Tyr Ser Tyr	CAA CAC AAN CTT ATA ACC TTA ACC CTA CTC TTT GGC TTC ACC GTT Gln His Xxx Leu Ile Thr Leu Thr Leu Leu Phe Gly Phe Thr Val	GGT TCG GTT CTC TAC TTC GTA ANC CGA CCC AAA CCG GTT TAC CTC Gly Ser Val Leu Tyr Phe Val Xxx Arg Pro Lys Pro Val Tyr Leu	GAC TAC TCC TGC TAC CTT CCA CCA CAA CAT CTT AGC GCT GGT ATC ASP Tyr Ser Cys Tyr Leu Pro Pro Gln His Leu Ser Ala Gly Ile	AAG ACC ATG GAA ATC TTT TAT CAA ATA AGA AAA TCT GAT CCT TTA Lys Thr Met Glu Ile Phe Tyr Gln Ile Arg Lys Ser Asp Pro Leu
CA	AAC Asn	AAA Lys	CTC	TTT Phe	GTT Val	TCT Ser

383	431	479	527	575	623	671
AAG Lys	GGA Gly	GAG Glu	AAC Asn	ATG Met	AAG Lys	AGT Ser
AGA Arg	GAG Glu	GAA Glu	AAC Asn	AGC	TCC	TGC
TTG	CCC	CGT Arg	GAG Glu	TCA Ser	ACT Thr	GGT Gly
TTC Phe	GGC Gly	GCG Ala	TTC	AAC Asn	AAT Asn	ATG
GAT Asp	TAC	TCG	CTA	GTG Val	GTT Val	GGA Gly
CTT Leu	ACC	GCG Ala	AAT Asn	GTG Val	GTA Val	GGA
TCT Ser	GAA Glu	TTA Leu	AAA Lys	CTT Leu	ATG Met	CTT
TCT Ser	GAT Asp	AAT Asn	CTA Leu	ATA Ile	GCG Ala	AAT Asn
TCG	GGC Gly	AAG Lys	GCG Ala	GGT G1Y	TCC	TTT Phe
GAT Asp	CTA Leu	AGG Arg	$_{\rm GGT}$	ATT Ile	TTA Leu	AGC Ser
GAT Asp	GGT G1y	CCG	AAC Asn	GAG Glu	TCG	AAA Lys
TTA Leu	TCA Ser	CCT	ATC Ile	AAA Lys	CCT	ATC Ile
GCA Ala	CGT Arg	ATT Ile	GTA Val	CCT	ACT	AAC Asn
GTG Val	GAG	GAG Glu	CAA Gln	AAC Asn	CCG	AGC
AAC Asn	CAA Gln	TTT Phe	GAG	GTT Val	AAT Asn	CGA Arg
CGA Arg	ATT Ile	CTG	ACG	AAA Lys	TTT	CTC

TG. 13E

719	767	815	863	911	949	1007
CAT His	AAC Asn	TTC	CGA Arg	GGA	AGC	GGG G1y
GTT	CAA	TTG	GAT	ACC	GAT	GCC
Val	Gln		Asp	Thr	Asp	Ala
CAT	ACT	TGC	666	CAT	GAT	GTT
His	Thr		G1y	His	Asp	Val
TTG	ATC	AAT	CCG	ACG	GAA	GTT
	Ile	Asn	Pro	Thr	Glu	Val
TTG	AAC Asn	TCG Ser	AAG Lys	CGA	GAA Glu	ACC Thr
GAC	GAG Glu	GTT Val	AAC Asn	GTT Val	CAA	ATA Ile
AAA	ACA	ATG	TCC	ACG	CGG	GAC
Lys	Thr	Met	Ser	Thr	Arg	Asp
GCT	AGC	ATG	CTC	CAC	GTG	AAA
Ala	Ser	Met		His	Val	Lys
CTA Leu	GTG Val	TCC Ser	CTG	GCT Ala	TGT Cys	TCA
GAT Asp	GTG Val	AGA Arg	ATT Ile	CTA	GGA G1y	TTG
ATT	CTT	AAC	GCG	AAG	TTT	AGT
Ile	Leu	Asn	Ala	Lys	Phe	
GCC	GCT	GAT	GCA	TAC	TCT	GTT
Ala	Ala	Asp	Ala		Ser	Val
ATC	\mathtt{TAT}	GGT	GGG	AAG	AAG	GGA
Ile		G1y	Gly	Lys	Lys	Gly
GTT	ACA	ACC	GGT	TCC	GAC	ACC
Val		Thr	Gly	Ser	Asp	Thr
GGT	AAC	TAT	GTC	CGG	GAC	AAA
Gly	Asn	Tyr	Val	Arg	ASP	Lys
GCT	AAA	ATT	CGT	AGA	GCT	GGT
Ala	Lys	Ile	Arg	Arg	Ala	Gly

FIG. 13(

1055	1103	1151	1199	1247	1295	1343
CCT	CTA	GCA	GTG	AGA	TAT	AAA
	Leu	Ala	Val	Arg	Tyr	Lys
CTT	AAA	CTT	GAT	TCA	TGG	AAT
Leu	Lys	Leu	Asp	Ser		Asn
GTT	AAG	AAA	ATA	GCA	ATT	GGT .
Val	Lys	Lys	Ile	Ala	Ile	
TTG	GCC	TTC	GTG	GAG	TCA	AAA
Leu	Ala	Phe	Val	Glu		Lys
CCG Pro	GTA Val	GAT Asp	GCC Ala	GTG	AGT	AAG
GGT Gly	TTC	CCG Pro	AGA Arg	GAT ASD	TCT	ATG Met
TTG	ACA Thr	GTG Val	GGT G1y	ATA Ile	TCG	AGG
ACA	GTT	\mathtt{TAC}	GGA	CCG	ACA	GGA
Thr	Val		G1y	Pro	Thr	G1y
ACA	GTC	TAT	GCG	TCG	AAT	AAA
Thr	Val	Tyr	Ala		Asn	Lys
ATA	TTT Phe	CAC His	CAT His	CTA Leu	$^{666}_{61y}$	CCA Pro
AAC	CTT Leu	AAA Lys	ATT Ile	$^{\rm GGG}_{\rm G1Y}$	TTT Phe	GAG Glu
AAA	ATC	ATC	TGT	TTA	AGA	ATA
Lys	Ile	Ile	Cys	Leu	Arg	Ile
CAG Gln	AAA Lys	AAG Lys	TTC	AAC	CAT His	TAC
GTT	GAA	GAT	CAT	AAG	TTA	GCA
Val	Glu	ASP	His	Lys	Leu	Ala
ACG	AGC	AAA Lys	GAT Asp	GAG Glu	ACA Thr	TTA Leu
ATA	CTG	TTA	GTA	TTA	TCA	GAA
Ile		Leu	Val	Leu	Ser	Glu

1391	1439	1487	1537	1597	1657	1664
AAG TGT AAT AGT GCG GTT Lys Cys Asn Ser Ala Val	GCT AAT AGT CCT TGG GAA Ala Asn Ser Pro Trp Glu	TAT TCT GGT TCA TCA AAG Tyr Ser Gly Ser Ser Lys	TAATTTATGT ATCTCAAATG	ATGITGICCA CITICICITY TITITITICI TITITIAGIT ATAAITTAAI GGITACGAIG	GTT ACTAGTATAA AAAAAAAAA 1657	
GGT TTT Gly Phe	GCT TCA Ala Ser	CAA ATG Gln Met	CGG TCC Arg Ser	IYIYIYIA	CATGGGTGTT	
GGG TCA GG Gly Ser G	GTC GAG G Val Glu A	CCG GTT C2 Pro Val G	AAC GGT CC Asn Gly A	YTTTTT 1	AAGAATA (
ATA GCT GGT C Ile Ala Gly C	TTA CGC AAT G Leu Arg Asn V	CAC AAA TAT C His Lys Tyr F	CCT GTC CAA P	TCTCTTT TT	GTCGTTATAA ATAAAGAATA	
GCT TGC CAA A1 Ala Cys Gln I1	TGG GTC GCT TT Trp Val Ala Le	CAT TGC ATT CA His Cys Ile Hi	TCA GAG ACT CC Ser Glu Thr Pr	ATGTTGTCCA CTT	TTTTGTCTAG GTC	AAAAAA

(1G. 13E

51	66	147	195	243	291	339
CCCCAACA ATG ACC CAT AAC CAA AAC CAA CCT CAC CGG GCA Met Thr His Asn Gln Asn Gln Pro His Arg Ala	CAC GTT ACA AAC TCC GAT CAA AAC CAA AAC CAA AAC CAA His Val Thr Asn Ser Asp Gln Asn Gln Asn Gln	CCA AAT TTT CTC TTA TCT GTT CGG CTC AAA TAT GTA AAA 1. Pro Asn Phe Leu Leu Ser Val Arg Leu Lys Tyr Val Lys	CAT TAC CTA ATC TCC AAC GGT CTC TAC ATC CTC CTC CTC 19 His Tyr Leu Ile Ser Asn Gly Leu Tyr Ile Leu Leu Leu	GGC GGC ACA ATC GTA AAA CTC TCT TCC TTC ACA CTC AAC Gly Gly Thr Ile Val Lys Leu Ser Ser Phe Thr Leu Asn	CTC CTC TAC AAC CAC CTC CGT TTT CAT TTC CTC TCC GCC 29 Leu Leu Tyr Asn His Leu Arg Phe His Phe Leu Ser Ala	ACC GGA CTC TTA ATC TCT CTC TCC ACC GCC TAC TTC ACC TAR Thr Gly Leu Leu Ile Ser Leu Ser Thr Ala Tyr Phe Thr
	SG GTT CO Val	AT CTC sn Leu	G TAC Y Tyr	ic CTC	C TCT u Ser	GCT Ala
CTTTCTTCTT	CCG	AAT Asn	GGG G1y	CTC	CTC	CTC
CŢ	GTT Val	AAC	CTT	CCT	GAA Glu	ACA

'IG. 14

387	435	483	531	579	627	675	723
CCT	CAA Gln	ATC	CTT Leu	GCA Ala	$_{\rm G1y}^{\rm GGT}$	TTT	CTT Leu
AAA Lys	TCT	AAG Lys	GCT Ala	GAA Glu	ACC	TTG Leu	AAG Lys
TAC	CGA Arg	CAA Gln	GAA Glu	AAA Lys	AAA Lys	AGT	TAT
TGC	GAC	CAA Gln	CCT Pro	AGA Arg	GAG	TGT	AAG '
TCA Ser	ATG Met	TTT Phe	TTC	GCG Ala	CTT Leu	AAT Asn	AAT
TTC Phe	TTC	GCT Ala	TAC Tyr	GAA Glu	GTT Val	GTG Val	GTG Val
GAC Asp	ACA Thr	CTC	ACT	GAA Glu	TCT Ser	GTC Val	ATT I
CTC	GAA Glu	AAC Asn	AAA Lys	ATG Met	GAC Asp	CTT	ATG
CTC	CGT	GAC Asp	CAG Gln	TGT Cys	ATA Ile	ATC Ile	GCC Ala 1
TTC	ACT	GAA Glu	666 61y	CCT	GCT Ala	GGA Gly	TCC
GTC Val	TGC	ACA Thr	CTT	AAT Asn	GGA Gly	ATC Ile	CTT
CAT His	ATA Ile	TTC	GGT Gly	CCC	TTC	gat Asp	TCA Ser
CGT Arg	TTA Leu	ATC Ile	TCC	CCT	ATG Met	AAA Lys	CCG Pro
CCT	TCC	GGT Gly	AGA Arg	GTT Val	GTT Val	CCT Pro	ACG Thr
CGT	CCT Pro	GTA Val	GAA Glu	CGT Arg	ACT Thr	AAA Lys	CCG
ACC	GAC Asp	CGT	CTC	CTT	GAG Glu	GTG	AAT

771	819	867	915	963	1011	1059
GCT Ala	CCA	TGG	CGT Arg	ACC	GCC	GAA Glu
AGT Ser	CAA G1n	AAC Asn	TTC Phe	CGC Arg	GGA G1y	AAC Asn
TGT Cys	GTC Val	TTA Leu	ATC Ile	GAT	AAA (Lys (AAC A
GGT G1y	CAG Gln	ACC	TGC Cys	TCC	CAC His	GAC ASP
ATG Met	CTT Leu	ATA Ile	AAC Asn	TCC	ACC	GAA Glu
GGA Gly	CTT Leu	AAC Asn	TCT Ser	CGT	CGT Arg	CGA
GGT Gly	CAG Gln	GAG Glu	CTC	AAC Asn	GTC Val	CAA
CTC	AAA Lys	ACA Thr	CTT	TCA	CCC	TAC
AAT Asn	GCT Ala	AGC Ser	ATG Met	CTC	CAC His	GTT Val
$\mathtt{TAT}\\\mathtt{TYT}$	CTC	GTG Val	TCA	CTT Leu	ATC Ile	TGC
AGC Ser	GAT Asp	GTG Val	CGA Arg	GTA Val	CTC	GGC
TTG	ATT Ile	CTA	GAC Asp	GCC Ala	CAG Gln	TTT
ATT Ile	TCC Ser	GCA Ala	AAC Asn	GCC Ala	TAT Tyr	GCA Ala
AAC Asn	ATC Ile	TAC	$ ext{GGC}$	GGA Gly	AAA Lys	AAC
$_{\rm GGA}$	CTT Leu	TCA Ser	TTA Leu	GGA Gly	TCA	GAC
AGA Arg	GGA Gly	AAC Asn	TAC	ATG Met	CGT	AAC

1107	1155	1203	1251	1299	1347	1395
GCA Ala	CTA	GCT	TTC	GTG Val	GAA Glu	TCA
ATG Met	CCA Pro	GTG Val	GAT Asp	GCA Ala	ATG Met	AGC Ser
CTA	GGA Gly	CTC	CCC	AGA Arg	CAT His	AGT Ser
AAC Asn	CIC	ACA Thr	ATA Ile	GGT Gly	TGG	TCG
AAA Lys	ACA	CCA Pro	TAC Tyr.	GGA Gly	GAG Glu	ACT Thr
TCT Ser	ACA Thr	TTC	CCT	GCG Ala	TCA Ser	AAT Asn
CTC	ATA Ile	TTT Phe	AAG Lys	CAT His	TTA Leu	GGT Gly
TCA Ser	AAC Asn	CTG	ATA Ile	ATC Ile	GAT Asp	TTT Phe
GTC Val	ACA Thr	ATT Ile	AAA Lys	TGC	TTG Leu	CGG
GGA G1y	AAG Lys	CAG Gln	AAG Lys	TTC Phe	AAT Asn	AAC Asn
ATC Ile	CTC	GAA Glu	GTC Val	CAT His	AAG Lys	TTA
AAA Lys	GCT Ala	TCC Ser	AAA Lys	GAG Glu	GAG Glu	ACT
GCC	GAA Glu	ATG Met	TTC	TTC Phe	ATA Ile	ATG Met
ACC Thr	GGA Gly	CCA	ATC Ile	GCT Ala	GAG Glu	AGG
GAA Glu	GCC Ala	TTA Leu	AAA Lys	CTA	GAT Asp	TCG
GAA Glu	ATA Ile	GTC Val	CGA Arg	AAG Lys	CTT Leu	CCA Pro

'IG. 14D

	1701	TAATATITIGT TATGGTTITG TTCTTACGTA CGTACTITTAA GTGATTTAGT CTAAAAATAA 1701	TA
	1641	ATC ACT CCA GTT ACT TCT AAC TAGTGTTTTT TTTTTGGGTC CAACTAGGGA Ile Thr Pro Val Thr Ser Asn	AGG
	1587	AAT CCA TGG AGT GAG ATT CAT GAG TTT CCA GTT TCT GTT CCT Asn Pro Trp Ser Asp Glu Ile His Glu Phe Pro Val Ser Val Pro	AAG Lys
	1539	GCG GTT TGG AAA GCT TTG AGA ACC ATT GAT CCT ATT GAT GAG The Ala Val Trp Lys Ala Leu Arg Thr Ile Asp Pro Ile Asp Glu Lys	AGT
	1491	GAT AGG ACT TGC CAA ATT GCG TTT GGA TCG GGA TTT AAG TGT AAT ASP Arg Thr Cys Gln Ile Ala Phe Gly Ser Gly Phe Lys Cys Asn	GGA Gly
_	1443	r TGG TAT GAA CTT GCG TAT AGT GAA GCT AAA GGG AGG ATT AAG AGA 1 Trp Tyr Glu Leu Ala Tyr Ser Glu Ala Lys Gly Arg Ile Lys Arg	CTT

3. 14E

ATTGGTTTCA TAAAAAAA AAAAAAAAA A

48	96	144	192	240	288	336
TTG	TTG	TTA Leu	CTG	TGT Cys	GAG Glu	CAA Gln
TAT Tyr	ACG Thr	AAT Asn	ACT Thr	TCT Ser	TAT	TTC Phe
ATG Met	TCC	TTC Phe	$^{\rm GGG}_{\rm G1Y}$	TTC	TTC Phe	ACT
GCC Ala	CTC	AAG Lys	TTA Leu	GAT Asp	ATT Ile	TTA
AAC Asn	CAT His	CTT Leu	TTT Phe	GTG Val	GAG Glu	AAT
TCC	GCT Ala	CAG Gln	GTG Val	TTG	AGA	GAT
ATC	TTT Phe	GAA Glu	ATG Met	TAC	ACG	GAT
TTG	GCC Ala	TGG	CTT Leu	ATT Ile	TGC Cys	ACC
TAC	GTA Val	CTT Leu	AGC Ser	AAG Lys	ATA Ile	TTT Phe
CAT His	GCA Ala	CAT	TCG Ser	ACG Thr	CGT Arg	AAT Asn
TAC	CTA Leu	GTT Val	TGC Cys	CCG	GAG Glu	GGG G1y
GTA Val	CTT Leu	CTG	CTC	CGA	AAA Lys	ACT Thr
CTA	CCG	GAT Asp	ACT Thr	AGC	GAA Glu	CTA Leu
AAA Lys	GTG Val	CAA Gln	GTA Val	ATG Met	CCG	AAA Lys
CTT Leu	ATG Met	ATT Ile	TCA	TTC	AAG Lys	TCG
AAG Lys	TTA	ACG	CTG	TAT Tyr	TAC	AGA Arg

G. 15A

384	432	480	528	576	622
CCT	AGA Arg	GAG Glu	TGC Cys	CGG	ט
TTA Leu	GCT Ala	TTG	AAT Asn	AAT Asn	ATG
TAC	GAG Glu	TTG	GTG Val	GTT Val	666 G1y
ACG	GCG Ala	GAA Glu	GTG Val	GTG Val	66C 61y
AAC Asn	ATG Met	GAT Asp	CTT Leu	ATG Met	CTT
CAG Gln	TGT Cys	ATC Ile	ATT Ile	GCA	AAC
GGT G1y	CCG	GCG	$_{\rm GLy}^{\rm GGT}$	TCC	TAT
TTA	AAT Asn	GGT Gly	ATC Ile	CTG	AGT Ser
GGA G1y	CCC	TTC	GAT Asp	TCT Ser	ATA Ile
TCT	CCG	ATG Met	AAG Lys	CCG	ATC Ile
AGA	GTT Val	GTT Val	CCT	ACG	AAT Asn
GAA	CGG	ATG Met	AAA Lys	CCG	GGG G1v
ATC Ile	CTA	GAG	GTT Val	AAT Asn	AGA Arq
ATT	GTT Val	GCT	GGG G1y	TTC	CTT Leu
AAA Lys	GCC	GAG	ACC	TTG	AAG Lvs
AAG Lys	GAG Glu	AAG Lys	AAA Lys	AGC	TAC